## GMRES-based Iterative Refinement for Solving Linear Systems

Shutong Wu<br/>\* Valerie Vreugdenhil † Gaofeng Zhou ‡  ${\rm April}\ 18,\ 2024$ 

<sup>\*</sup>wu867@mcmaster.ca

 $<sup>^\</sup>dagger vreug dev@mcmaster.ca$ 

<sup>&</sup>lt;sup>‡</sup>zhoug28@mcmaster.ca

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## 1 Introduction

GMRES (Generalized Minimal Residual Method) and its variants based on iterative refinement (IR) are advanced numerical techniques for solving linear systems, particularly useful for large or ill-conditioned matrices where direct methods are inefficient or impractical.

GMRES is an iterative method for solving non-symmetric linear systems. It seeks to minimize the residual over a Krylov subspace generated by the matrix and the residual vector. The main steps in a GMRES iteration include the Arnoldi process to construct an orthogonal basis for the Krylov subspace, updating the QR factorization of the resulting Hessenberg matrix, and solving the resulting least squares problem to update the solution[3].

$$Ax = b \tag{1}$$

Assume that **A** is a real nonsingular **N** by **N** matrix and b is a real vector. Given an initial guess  $x^0$  for the solution, the algorithm generates approximate solutions  $x^n$ ,  $n = 1, 2, ..., \mathbf{N}$  from the linear variety

$$x^0 + \mathbf{K}_n(\mathbf{A}, r^o) \tag{2}$$

minimizing the Euclidean norm of the residual,

$$\|\mathbf{b} - A\mathbf{x}^n\| = \min_{\mathbf{u} \in \mathbf{x}_0 + \mathcal{K}_n(A, r_0)} \|\mathbf{b} - A\mathbf{u}\|, \tag{3}$$

where  $r^0 = b - \mathbf{A}x^0$  is the initial residual and  $K_n(\mathbf{A}, r^0)$  is the n - th Krylov subspace generated by  $\mathbf{A}, r^0$ ,

$$K_n(A, r^0) = \text{span}\{r^0, Ar^0, \dots, A^{n-1}r^0\}.$$
 (4)

Clearly,  $r^n \in r^0 + AK_n(A, r^0)$ . We call  $\mathbf{AK}_n(\mathbf{A}, r^0)$  a Krylov residual subspace.

Such an approximation always exists and is unique, It can be computed in many different

ways.

Most methods for computing the approximation  $x^n$  satisfying (3) start by constructing an orthonormal basis, called an Arnoldi basis, for the Krylove subspace (4). The recurrence for the basis vectors can be written in matrix for as

$$AV_n = V_{n+1} H_{n+1,n} (5)$$

Where  $V_{n+1}$  is the N by (n+1) matrix with the orthonormal basis vectors  $v_1, V_2, \ldots, V_{n+1}$  as its columns and  $H_{n+1,n}$  is the (n+1) by n upper Hesenberg matrix of the orthogonalization (and normalization) coefficients, n < N. The initial vector  $\mathbf{s} = \frac{\mathbf{r}^0}{\rho}$ , where  $\rho = ||\mathbf{r}^0||$ . The approximate solution  $x^n$  is then taken to be of the form  $x^n = x^0 + V_n y^n$ , where  $y^n$  is cosen to minimize

$$\|\mathbf{b} - A\mathbf{x}^n\| = \|\mathbf{r}^0 - AV_n\mathbf{y}^n\| = \|V_{n+1}(\mathbf{e}_1 - H_{n+1,n}\mathbf{y}^n)\| = \|(\mathbf{e}_1 - H_{n+1,n}\mathbf{y}^n)\|.$$
 (6)

The problem of solving apporximately the original N-dimensional system Ax = b is thus transformed to the n-demensional least squares problem

$$\|\mathbf{e}_1 - H_{n+1,n}\mathbf{y}^n\| = \min_{\mathbf{y}} \|\mathbf{e}_1 - H_{n+1,n}\mathbf{y}\|, \quad \mathbf{x}^n = \mathbf{x}^0 + V_n\mathbf{y}^n.$$
 (7)

We call  $b - Ax^n$  a true residual,  $\rho e_1 - H_{n+1,n}y^n$  an Arnoldi residual. While the norms of these two vector are the same in exact arithmetic.

There are sereval algorihms for computing the Arnoldi basis, while GMRES has played a significant role since it has been invented.

GMRES-IR extends GMRES by incorporating mixed precision strategies to improve computational efficiency and robustness. GMRES-IR operates by using different precision levels throughout the computation process. It typically involves a sequence where a lower precision

is used for certain operations to save computational cost, while higher precision is reserved for critical steps to ensure the accuracy of the solution. This method is particularly effective when solving nearly singular or ill-conditioned systems where high precision is required to achieve an accurate solution .

# 1.1 Key Concepts of GMRES and IR algorithms based on GM-RES

#### • Multiple Precision Levels:

- GMRES-IR strategically uses different precision levels through the computation process. Lower precision is used where high accuracy is less crucial, reducing computational cost and memory usage.
- Higher precision is used in critical steps, particularly in the evaluation of residuals
  and the final solution steps, where maintaining numerical accuracy is paramount.
  This approach helps in managing the trade-off between computational speed and
  accuracy effectively.

#### • Iterative Refinement:

- In GMRES-IR, the initial solution to the linear system is typically computed using a standard GMRES iteration in a lower precision format (e.g., single precision).
   This serves to obtain a coarse solution quickly.
- Once this solution is computed, the residual (the difference between the left-hand side and right-hand side of the linear equation using the current solution) is recalculated in higher precision (e.g., double precision). This step is crucial because it allows the detection of small errors that lower precision may miss.

The refined solution is then improved iteratively by applying additional GMRES cycles targeted at the residual, computed in higher precision. This process can be repeated multiple times to progressively enhance the solution accuracy.

#### • Preconditioning:

- Preconditioning in GMRES-IR is vital for improving the convergence rates of the iterative process. A preconditioner transforms the original system into one that is easier (i.e., faster) to solve iteratively.
- The preconditioner itself can be computed in a lower precision to save computational resources, and then applied during the higher precision solution phase.
   This is effective in reducing overall computational times while still maintaining a robust approach towards convergence.

#### • Application-Specific Adjustments:

The configuration of precision levels, the choice of preconditioner, and the number of refinement iterations can be tailored based on specific application needs and hardware capabilities. This customization allows GMRES-IR to be optimized for specific scenarios, such as sparse systems, systems arising in fluid dynamics, or systems that require high levels of precision due to their sensitivity.

#### 1.2 Comparative Analysis of GMRES-IR and LU-IR

This section conducts a comparative analysis of GMRES-based IR(GMRES-IR) and LU-based IR(LU-IR) in terms of accuracy, convergence, and performance in a mixed-precision setting.

#### 1.2.1 LU-IR

Iterative refinement has traditionally used LU factorization as the inner solver. A basic iterative refinement algorithm using LU factorization can be described as follows:

**Algorithm 1** LU-IR.  $A \in \mathbb{R}^{n \times n}$  is nonsingular and  $b \in \mathbb{R}^n$ . Three precisions are used, satisfying  $u_r \leq u \leq u_l$ .

- 1: Compute the factorization A = LU in precision  $u_l$ .
- 2: Solve  $LUx_1 = b$  by substitution in precision  $u_l$ .
- 3: for  $i = 1 : i_{\text{max}}$  or until converged do
- 4: Compute  $r_i = b Ax_i$  in precision  $u_r$ .
- 5: Solve  $LUd_i = r_i$  by substitution in precision  $u_l$ .
- 6: Update  $x_{i+1} = x_i + d_i$  in precision u.
- 7: end for

Here the most computational heavy step is the LU factorization, which costs  $O(n^3)$  flop. The substitutions on line 5 costs in total  $O(Kn^2)$  flops, which is negligible compared to LU factorization for large n and reasonable K (K is the total number of iterations). So by adopting a lower precision for  $u_l$ , the computational cost can be greatly reduced. With half-precision availability ( $u_{16}$ ), the potential speedup with  $u = u_r = u_{64}$  and  $u_l = u_{16}$  on an NVIDIA P100 GPU can reach 2.7.

The convergence of LU-IR depends on various factors, including numerical stability of the LU decomposition, precision scheme, and also the condition of A. Specifically, with  $\kappa(A)u_l \ll 1$  and a numerically stable LU decomposition, convergence is guaranteed. However, when lower precision is applied (especially in half-precision schemes), the requirement for  $\kappa(A)$  tightens. For instance, with fp16, the unit round-off  $u_{16} = u_l \approx 2^{-11}$ , rendering  $\kappa(A) \ll 2000$ .

With bfloat16, the unit round-off  $u_{16} = u_l \approx 2^{-8}$ , giving an even tighter bound  $\kappa(A) \ll 300$ . This marks one limitation of LU-based IR, which is the convergence heavily relies on the system condition and unit round-off. Especially in a half-precision computing setting, this condition limitation of LU-IR render itself a strict and picky solver[1].

The above algorithm achieves a relative error bounded approximately by  $3n||A^{-1}||\hat{L}||\hat{U}||u_l$ .

#### 1.2.2 **GMRES-IR**

The GMRES-based iterative refinement (GMRES-IR) in a mixed-precision setting can be described as follows:

**Algorithm 2** GMRES-IR.  $A \in \mathbb{R}^{n \times n}$  is nonsingular and  $b \in \mathbb{R}^n$ . Five precisions are used:  $u_r, u_g, u_p, u, u_l$ .

- 1: Compute the factorization A = LU in precision  $u_l$ .
- 2: Solve  $LUx_1 = b$  by substitution in precision  $u_l$ .
- 3: for  $i = 1 : i_{\text{max}}$  or until converged do
- 4: Compute  $r_i = b Ax_i$  in precision  $u_r$ .
- 5: Solve  $U^{-1}L^{-1}Ad_i = U^{-1}L^{-1}r_i$  by GMRES in precision  $u_g$ , performing the products with  $U^{-1}L^{-1}A$  in precision  $u_p$ .
- 6: Compute  $x_{i+1} = x_i + d_i$  in precision u.
- 7: end for

An important feature of GMRES-IR is its applicability to extremely ill-condition matrices.

In algorithm 2 line 4, denote the relative error:

$$\xi_i = \frac{\|d_i - \hat{d}_i\|_{\infty}}{\|d_i\|_{\infty}}$$

Let

$$\mu_i = \frac{\|A(\mathbf{x}_i - \hat{\mathbf{x}}_i)\|_{\infty}}{\|A\|_{\infty} \|\mathbf{x}_1 - \hat{\mathbf{x}}_i\|_{\infty}} \le 1,$$

$$\phi_i = 2 \min(\operatorname{cond}(A), \kappa_{\infty}(A)\mu_i)u_l + \xi_i,$$

where cond(A) =  $|||A^{-1}||A||x|||_{\infty}$ . As long as  $\phi_i$  is sufficiently less than 1,  $\xi_i$  is reduced at each iteration until an iterate  $\hat{x}$  is obtained such that:

$$\frac{\|x - \hat{x}\|_{\infty}}{\|x\|_{\infty}} \lesssim u + 4p \ cond(A, x)u_r$$

where p is the maximum number of nonzeros in any row of  $[A \ b]$ , and cond(A,x) is defined as

$$cond(A, x) = \frac{\||A^{-1}||A||x|\|_{\infty}}{\|x\|_{\infty}}.$$

This accuracy bound depends on precisions u and  $u_r$ , and is independent of precision  $u_l$  and  $x_1$ . This motivates adopting a higher precision for the residual  $u_r$ . It's experimented that by adopting  $u = u_l$  and  $u_r = u_g = u_p = u^2$ ,  $\xi_i \approx u$  as long as  $\kappa(A) \approx u^{-1}$ , indicating that the error is of the same order as the precision of the computation regardless of the system condition. This is a significant result, as it suggests that systems that are ill-conditioned, which would typically be problematic due to their high sensitivity to errors, can still be solved to a high degree of accuracy using GMRES-IR, with the caveat that a higher precision is used in part of the computation to control the growth of errors. This demonstrates the advantage of GMRES-based IR over LU-based IR, where the requirement for system condition is greatly relaxed with reasonable computational overhead[2].

In terms of error behavior, one caveat of using GMRES-based method is the possibility of error accumulation inside the subspace iteration. As a Krylov subspace iterative method, round-off errors accumulated along the recurrent chain might result in a considerable deviation from the true value. Here the trade-off is, the longer the Krylov subspace constructed, the more dimensions available and therefore higher accuracy solution and faster convergence, while at the same time it also leads to larger chance of round-off error accumulation as a result of long inner iteration chain. A possible mitigation to such problem is the constant restart, where the Krylov subspace is reset after a preset amount of iterations. In the itera-

tive refinement setting, as GMRES is employeed as a inner solver, the returning to the outer loop itself acts as a reset on the subspace. Thus the enforced restart is more likely to be triggered with a low or no preconditioning, as in such case more calculation is carried out inside the Krylov subspace.

In the above algorithm, the preconditioner  $U^{-1}L^{-1}$  is used, which costs  $O(n^3)$  flops. To avoid such cost, it's also possible to use a less computational expensive preconditioner for GMRES-IR, or even no preconditioner at all. With this tradeoff, larger iteration count is expected, but convergence is still guaranteed with condition  $\kappa(A)u_l \ll 1$  met. Examples of less costy preconditioners used in GMRES-IR include incomplete LU factorization, block Jacobi and polynomial preconditioners[4], With less restrict preconditioning and system condition requirements, GMRES-IR can choose more flexible precision schemes compared to LU-IR, thus utilizing mixed-precision computing to a higher level.

## 2 Experiments

#### 2.1 Matrix Generation and Results

We use several functions to generate those matrices we need to do this experiment.

```
function A = genMatrix(size, cond)

A = 2*rand(size);

[u, s, v] = svd(A);

s = diag(s);

s = s(1)*( 1-((cond-1)/cond)*(s(1)-s)/(s(1)-s(end)));

s = diag(s);

A = u*s*v';

end
```

Listing 1: generating matrix

Small size matrix:

```
cond = [1e2, 1e3, 1e4, 1e5, 1e6, 1e7, 1e8, 1e9];
matrices = cell(size(cond,2));
for i=1:size(cond,2)

matrices{i} = genMatrix(500, cond(i));
end

relresgm = zeros(size(matrices,2),1);
relreslu = zeros(size(matrices,2),1);
itergm = zeros(size(matrices,2),1);
iterlu = zeros(size(matrices,2),1);
timegm = zeros(size(matrices,2),1);
timegm = zeros(size(matrices,2),1);
timelu = zeros(size(matrices,2),1);

timelu = zeros(size(matrices,2),1);

for i = 1:size(matrices,2)
b = rand(size(matrices,2),1);
```

```
tic
16
      [x, flag, relres, iter] = gmres(matrices{i}, b, [], 1e-6, 30);
17
      timegm(i) = toc;
18
      relresgm(i) = relres;
19
      flags(i) = flag;
20
      itergm(i) = iter(2);
21
22
      tic
      [x, relres, iter] = iterref(matrices{i}, b);
24
      timelu(i) = toc;
      relreslu(i) = relres(size(relres,2));
26
      iterlu(i) = iter;
28 end
30 figure (1)
31 semilogy(cond, relreslu);
32 title("luir: accuracy vs condition (small matrix - 500x500)")
saveas(gcf,'luir_acc_smat.png')
34 figure (2)
semilogy(cond, relresgm);
36 title("gmres: accuracy vs condition (small matrix - 500x500)")
saveas(gcf,'gmres_acc_smat.png')
39 figure (3)
40 plot(cond, iterlu);
41 title("luir: iterations vs condition (small matrix - 500x500)")
saveas(gcf,'luir_iter_smat.png')
43 figure (4)
44 plot(cond, itergm);
45 title("gmres: iterations vs condition (small matrix - 500x500)")
saveas(gcf,'gmres_iter_smat.png')
```

```
figure(5)

figure(5)

plot(sizes, timelu);

title("luir: time vs matrix size (low condition number)")

saveas(gcf,'luir_acc_lcond.png')

figure(6)

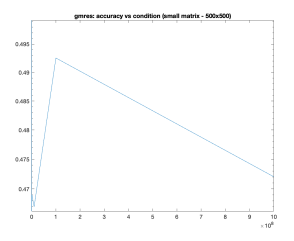
plot(sizes, timegm);

title("gmres: time vs matrix size (low condition number)")

saveas(gcf,'luir_acc_lcond.png')
```

Listing 2: Small size mtrix

With this small matrix, we got the results as bellow:



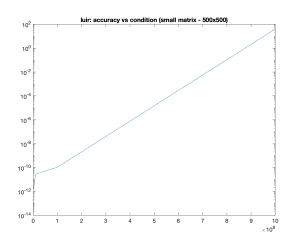


Figure 1: Accuracy with condition number with GMRES

Figure 2: Accuracy with condition number with LU-IR.

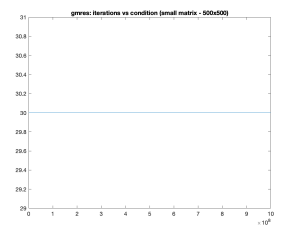
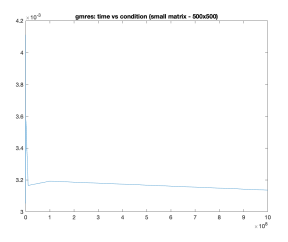


Figure 3: Iterations with condition number with  $\operatorname{GMRES}$ 

Figure 4: Iterations with condition number with LU-IR.



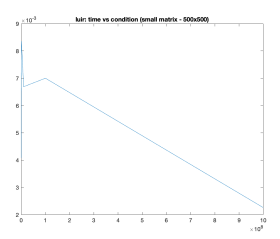


Figure 5: Time with condition number with GMRES

Figure 6: Time with condition number with LU-IR.

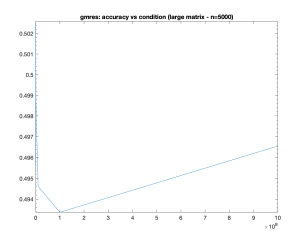
Large size matrix:

```
cond = [1e2, 1e3, 1e4, 1e5, 1e6, 1e7, 1e8, 1e9];
2 matrices = cell(size(cond,2));
3 for i=1:size(cond,2)
      matrices{i} = genMatrix(5000, cond(i));
5 end
7 relresgm = zeros(size(matrices,2),1);
8 relreslu = zeros(size(matrices,2),1);
9 itergm = zeros(size(matrices,2),1);
iterlu = zeros(size(matrices,2),1);
timegm = zeros(size(matrices,2),1);
timelu = zeros(size(matrices,2),1);
flags = zeros(size(matrices,2),1);
for i = 1:size(matrices,2)
      b = rand(size(matrices{i},1),1);
      tic
17
      [x, flag, relres, iter] = gmres(matrices{i}, b, [], 1e-6, 30);
18
      timegm(i) = toc;
19
20
      relresgm(i) = relres;
      flags(i) = flag;
21
      itergm(i) = iter(2);
22
23
      tic
      [x, relres, iter] = iterref(matrices{i}, b);
25
      timelu(i) = toc;
26
      relreslu(i) = relres(size(relres,2));
2.7
      iterlu(i) = iter;
29 end
30
```

```
31 figure (1)
32 semilogy(cond, relreslu);
33 title("luir: accuracy vs condition (large matrix - n=10000)")
saveas(gcf,'luir_acc_lmat.png')
35 figure(2)
semilogy(cond, relresgm);
37 title("gmres: accuracy vs condition (large matrix - n=10000)")
saveas(gcf,'gmres_acc_lmat.png')
40 figure (3)
41 plot(cond, iterlu);
42 title("luir: iterations vs condition (large matrix - n=10000)")
saveas(gcf,'luir_iter_lmat.png')
44 figure (4)
45 plot(cond, itergm);
46 title("gmres: iterations vs condition (large matrix - n=10000)")
saveas(gcf,'gmres_iter_lmat.png')
49 figure (5)
50 plot(sizes, timelu);
51 title("luir: time vs matrix size (low condition number)")
52 saveas(gcf,'luir_acc_lcond.png')
53 figure(6)
54 plot(sizes, timegm);
55 title("gmres: time vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
```

Listing 3: Large size matrix

With large matrix as above, we got those results as bellow:



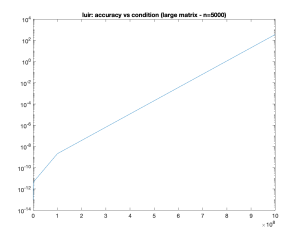


Figure 7: Accuracy with condition number with GMRES

Figure 8: Accuracy with condition number with LU-IR.

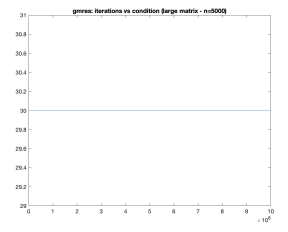


Figure 9: Iterations with condition number with GMRES

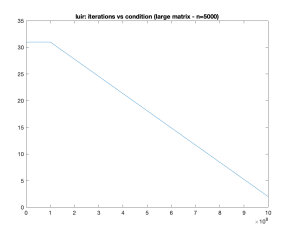
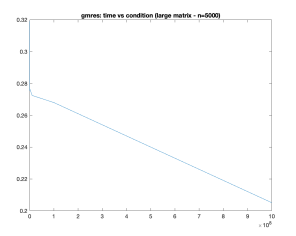


Figure 10: Iterations with condition number with LU-IR.



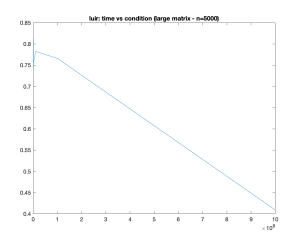


Figure 11: Time with condition number with  $\operatorname{GMRES}$ 

Figure 12: Time with condition number with LU-IR.

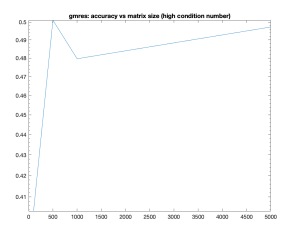
Matrix with high condition number:

```
sizes = [10,100,500,1000, 5000];
matrices = cell(size(sizes,2));
3 for i=1:size(sizes,2)
      matrices{i} = genMatrix(sizes(i), 1e9);
5 end
7 relresgm = zeros(size(matrices,2),1);
8 relreslu = zeros(size(matrices,2),1);
9 itergm = zeros(size(matrices,2),1);
iterlu = zeros(size(matrices,2),1);
timegm = zeros(size(matrices,2),1);
timelu = zeros(size(matrices,2),1);
flags = zeros(size(matrices,2),1);
for i = 1:size(matrices,2)
      b = rand(size(matrices{i},1),1);
      tic
17
      [x, flag, relres, iter] = gmres(matrices{i}, b, [], 1e-6, 30);
18
      timegm(i) = toc;
19
20
      relresgm(i) = relres;
      flags(i) = flag;
21
      itergm(i) = iter(2);
22
23
      tic
      [x, relres, iter] = iterref(matrices{i}, b);
25
      timelu(i) = toc;
26
      relreslu(i) = relres(size(relres,2));
2.7
      iterlu(i) = iter;
29 end
30
```

```
31 figure(1)
32 semilogy(sizes, relreslu);
33 title("luir: accuracy vs matrix size (high condition number)")
saveas(gcf,'luir_acc_hcond.png')
35 figure (2)
36 semilogy(sizes, relresgm);
137 title("gmres: accuracy vs matrix size (high condition number)")
saveas(gcf,'gmres_acc_hcond.png')
40 figure (3)
41 plot(sizes, iterlu);
42 title("luir: iterations vs matrix size (high condition number)")
saveas(gcf,'luir_iter_hcond.png')
44 figure (4)
45 plot(sizes, itergm);
46 title("gmres: iterations vs matrix size (high condition number)")
saveas(gcf,'luir_iter_hcond.png')
49 figure (5)
50 plot(sizes, timelu);
51 title("luir: time vs matrix size (low condition number)")
52 saveas(gcf,'luir_acc_lcond.png')
53 figure(6)
54 plot(sizes, timegm);
55 title("gmres: time vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
```

Listing 4: Matrix with high contional number

With matrix with high condition number, we got results as bellow:



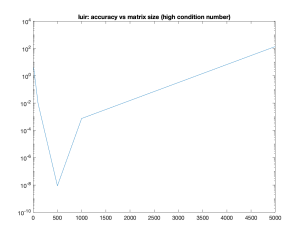
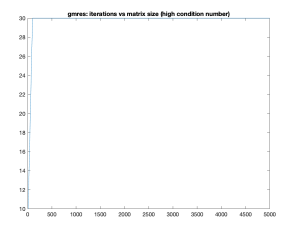


Figure 13: Accuracy with size of matrix with GMRES

Figure 14: Accuracy with size of matrix with LU-IR.



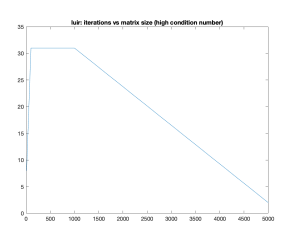
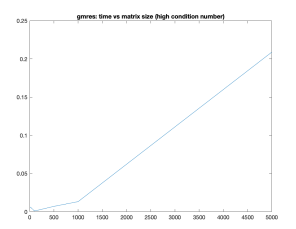


Figure 15: Iterations with size of matrix with  $\operatorname{GMRES}$ 

Figure 16: Iterations with size of matrix with LU-IR.



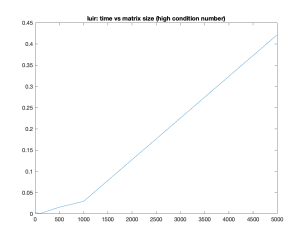


Figure 17: Time with size of matrix with GMRES

Figure 18: Time with size of matrix with LU-IR.

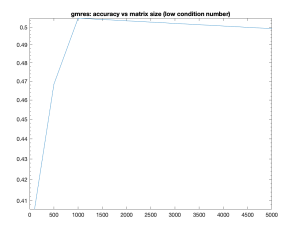
Matrix with low condition number:

```
sizes = [10,100,500,1000,5000];
matrices = cell(size(sizes,2));
3 for i=1:size(sizes,2)
      matrices{i} = genMatrix(sizes(i), 1e2);
5 end
7 relresgm = zeros(size(matrices,2),1);
8 relreslu = zeros(size(matrices,2),1);
9 itergm = zeros(size(matrices,2),1);
iterlu = zeros(size(matrices,2),1);
timegm = zeros(size(matrices,2),1);
timelu = zeros(size(matrices,2),1);
flags = zeros(size(matrices,2),1);
for i = 1:size(matrices,2)
      b = rand(size(matrices{i},1),1);
      tic
17
      [x, flag, relres, iter] = gmres(matrices{i}, b, [], 1e-6, 30);
18
      timegm(i) = toc;
19
20
      relresgm(i) = relres;
      flags(i) = flag;
21
      itergm(i) = iter(2);
22
23
      tic
      [x, relres, iter] = iterref(matrices{i}, b);
25
      timelu(i) = toc;
26
      relreslu(i) = relres(size(relres,2));
2.7
      iterlu(i) = iter;
29 end
30
```

```
31 figure(1)
32 semilogy(sizes, relreslu);
33 title("luir: accuracy vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
35 figure(2)
36 semilogy(sizes, relresgm);
37 title("gmres: accuracy vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
40 figure (3)
41 plot(sizes, iterlu);
42 title("luir: iterations vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
44 figure (4)
45 plot(sizes, itergm);
46 title("gmres: iterations vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
49 figure (5)
50 plot(sizes, timelu);
51 title("luir: time vs matrix size (low condition number)")
52 saveas(gcf,'luir_acc_lcond.png')
53 figure(6)
54 plot(sizes, timegm);
55 title("gmres: time vs matrix size (low condition number)")
saveas(gcf,'luir_acc_lcond.png')
```

Listing 5: Matrix with low conditional number

With Matrix with low condition number as above, we got results as bellow:



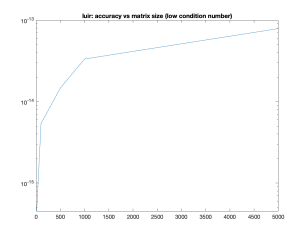
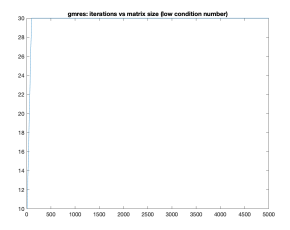


Figure 19: Accuracy with size of matrix with  $\operatorname{GMRES}$ 

Figure 20: Accuracy with size of matrix with LU-IR.



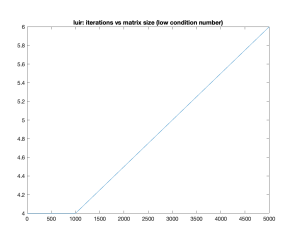


Figure 21: Iterations with size of matrix with  $\operatorname{GMRES}$ 

Figure 22: Iterations with size of matrix with LU-IR.

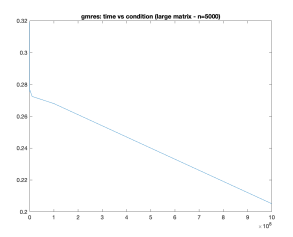


Figure 23: Time with size of matrix with GMRES

Figure 24: Time with size of matrix with LU-IR.

## 3 Results Analysis

#### 1. Small size matrix:

#### • GMRES:

- Time vs.Condition Number:
  - \* The time required by GMRES slightly decreases as the condition number increases, but it stabilizes for higher condition numbers.
  - \* This pattern might be due to the nature of iterative methods like GMRES that can exhibit complex relationships with condition numbers depending on factors such as preconditioning, stopping criteria, and the specific characteristics of the matrix.

#### - Iterations vs. Condition Number:

- \* GMRES shows a remarkably stable number of iterations across the range of condition numbers.
- \* This stability is a hallmark of well-implemented iterative methods that are not significantly impacted by the condition number, particularly for small matrix sizes.

#### - Accuracy vs. Condition Number:

\* GMRES's accuracy deteriorates as the condition number increases, which is typical behavior as higher condition numbers usually result in less accurate solutions due to the potential for error amplification.

#### • LU-IR:

- Time vs. Condition Number:
  - \* The graph exhibits a decreasing trend in time with increasing condition numbers, which is unconventional as higher condition numbers typically

imply a more ill-conditioned matrix and could require more computational effort.

\* A potential explanation for this trend could be the specific characteristics of the matrix at different condition numbers or the way the iterative refinement is implemented, possibly needing fewer corrections for higher condition numbers.

#### - Iterations vs. Condition Number:

- \* The downward trend in the number of iterations as the condition number increases aligns with the trend in computational time.
- \* This suggests that for LU-IR, higher condition numbers are somehow resulting in fewer iterations to reach convergence, which is an interesting observation and could be due to the interplay between the condition number and the convergence criteria of the algorithm.

#### - Accuracy vs. Condition Number:

- \* The accuracy graph shows a clear inverse correlation between accuracy and condition number; as the condition number increases, the accuracy of the solution decreases.
- \* This is expected since a higher condition number typically indicates a matrix that is closer to being singular or ill-conditioned, leading to larger errors in the solution.

#### • Comparative Analysis:

- Comparing the LU-IR to GMRES, GMRES shows better stability in terms of iterations, but both methods display the expected decrease in accuracy with higher condition numbers.
- For the LU-IR method, there's an interesting correlation where both the

time and number of iterations decrease with increasing condition numbers, yet the accuracy worsens. This might suggest that the iterative refinements are prematurely terminated or become less effective as the matrix becomes more ill-conditioned.

The computational time trends for both methods against the condition number are not typical and might suggest implementation details, such as the effect of stopping criteria or preconditioning, which have a more significant impact than the condition number itself.

#### • Conclusion:

The results suggest that both algorithms maintain a certain level of robustness in the face of varying condition numbers. However, the loss of accuracy with increasing condition number is consistent with the mathematical theory that higher condition numbers lead to more numerical instability.

#### 2. Large size matrix:

#### • GMRES:

- Time vs. Condition Number:
  - \* GMRES shows a mild decrease in time with increasing condition numbers, eventually reaching a stable point.
  - \* This might suggest that GMRES, as an iterative method, is less affected by higher condition numbers, possibly due to good preconditioning that mitigates the effects of ill-conditioning.

#### - Iterations vs. Condition Number:

\* The GMRES method displays a consistent number of iterations across various condition numbers, indicating stable convergence properties.

\* The stability of iterations suggests that GMRES is quite robust to changes in the condition number, at least for the matrix sizes and condition numbers tested.

#### - Accuracy vs. Condition Number:

\* GMRES accuracy decreases with an increase in the condition number, which is consistent with expectations. High condition numbers lead to lower accuracy due to the potential amplification of errors in iterative methods.

#### • LU-IR:

#### - Time vs. Condition Number:

- \* The computational time decreases with an increase in condition numbers, which is a trend that does not align with the typical expectation that higher condition numbers lead to more computational work.
- \* The decrease may be explained by certain optimizations or behaviors specific to the LU-IR method that make the computational cost less sensitive to changes in the condition number for this range of matrix sizes.

#### - Iterations vs. Condition Number:

- \* The decreasing number of iterations required as the condition number increases further supports the observed decrease in computational time.
- \* This suggests that the algorithm requires fewer iterations to converge for matrices with higher condition numbers. However, this is an unusual pattern and may warrant investigation into the specifics of the implementation or the nature of the matrices used.
- The initial overhead might include factors like memory allocation, construction of the LU decomposition, and other once-off computations.

#### - Accuracy vs. Condition Number:

\* As expected, the accuracy worsens with increasing condition numbers, reflecting the typical behavior where higher condition numbers indicate a system that is more difficult to solve accurately due to increased numerical instability.

#### • Comparative Analysis:

- Both methods exhibit the expected degradation of accuracy with higher condition numbers.
- The number of iterations and computational time patterns for LU-IR with respect to condition number are unconventional and could be specific to the particular matrices tested or the implementation of the algorithm.
- GMRES shows a mild decrease in time but maintains a consistent number of iterations, indicating its potential robustness and efficiency as the condition number increases.

#### • Conclusion:

- The observations suggest that GMRES is more stable with respect to the number of iterations and time against increasing condition numbers for the tested large matrix. This aligns with the expectations of iterative methods like GMRES, which can be designed to be more resilient to changes in condition numbers.
- For LU-IR, the decrease in time and iterations with increasing condition numbers is an intriguing behavior that may indicate unique properties of the method or the particular systems being solved.
- The degradation in accuracy for both methods is consistent with the mathematical challenges posed by high condition numbers.

#### 3. Low condition numbered matrix:

#### • GMRES:

- Time vs. Matrix Size:
  - \* The computational time for GMRES increases nearly linearly with matrix size, which is excellent scaling behavior for an iterative method, particularly since the theoretical complexity for solving linear systems using Krylov subspace methods like GMRES is often higher.

#### - Iterations vs. Matrix Size:

\* The number of iterations remains almost constant, which is indicative of the robustness of the GMRES algorithm with respect to matrix size, especially given that the condition number is low.

#### - Accuracy vs. Matrix Size:

\* GMRES maintains high accuracy across matrix sizes, showing a slight decrease as size increases. This is consistent with expectations as larger systems can amplify numerical errors, but the impact here is minimal, benefiting from the low condition number.

#### • LU-IR:

#### - Time vs. Matrix Size:

\* The time taken by LU-IR increases in what appears to be a quadratic manner with the size of the matrix, which is expected as the LU decomposition has a computational complexity of  $O(n^3)$  for dense matrices. However, the observed trend might suggest a slightly better performance, possibly due to the low condition number or specific matrix structure allowing faster computation.

#### - Iterations vs. Matrix Size :

\* The number of iterations increases linearly with the matrix size. Since the condition number is low, the linear systems are well-conditioned, and thus the iterative refinement process converges relatively quickly, leading to a predictable increase in iterations with size.

#### - Accuracy vs. Matrix Size:

\* The accuracy remains high across different matrix sizes, which suggests that the LU-IR method is capable of producing accurate results for well-conditioned matrices. The slight decrease in accuracy with size might be attributed to the accumulation of numerical errors as the size increases.

#### • Comparative Analysis:

- Both LU-IR and GMRES show good performance in terms of accuracy for well-conditioned matrices, even as the matrix size increases.
- GMRES demonstrates exceptionally stable and efficient scaling in terms of time with matrix size, with only a slight increase in iterations needed.
- The quadratic increase in computational time for LU-IR is typical for direct decomposition methods, while the near-linear scaling of GMRES highlights the advantages of iterative methods for large-scale computations.

#### • Conclusion:

- LU-IR shows expected scaling of time and iterations with matrix size but maintains high accuracy, which is a positive outcome for low condition numbers.
- GMRES exhibits remarkable stability in the number of iterations and maintains accuracy, making it an excellent choice for larger matrices even as they grow in size.
- The low condition number of the matrices aids in both algorithms' perfor-

mance, ensuring numerical stability and reliable accuracy.

#### 4. High condition numbered matrix:

#### • GMRES:

#### - Time vs. Matrix Size:

\* GMRES shows a near-linear increase in computational time with the size of the matrix. This performance is remarkable, considering that iterative methods can be sensitive to matrix conditioning. The near-linear relationship might be a consequence of effective preconditioning or the specific matrix structures enabling more efficient computation.

#### - Iterations vs. Matrix Size:

\* The number of iterations needed by GMRES also increases linearly with matrix size. This is typical for iterative methods, where the number of iterations needed tends to increase with problem size but is also influenced by the conditioning of the matrix and the effectiveness of any preconditioner used.

#### - Accuracy vs. Matrix Size:

\* The accuracy graph suggests that the solutions become less accurate as the matrix size increases. This is an expected trend for high condition number matrices, as the error is more likely to propagate and accumulate over iterations in large-scale problems.

#### • LU-IR:

#### - Time vs. Matrix Size:

\* The computational time for LU-IR grows almost linearly with the matrix size, which is better than the expected cubic time complexity of the LU

decomposition. This may indicate that the specific characteristics of the matrices being used allow for faster computation, even though they are poorly conditioned.

#### - Iterations vs. Matrix Size:

\* The graph shows a decreasing number of iterations required as the matrix size increases, which is counterintuitive because poorly conditioned matrices typically require more iterative refinement. This could be a result of the iterative refinement process reaching a specified precision limit sooner for larger matrices, or due to specific properties of the matrices being tested.

#### - Accuracy vs. Matrix Size:

\* The accuracy decreases as the matrix size increases, which aligns with expectations; larger and poorly conditioned matrices amplify numerical errors, leading to less accurate solutions.

#### • Comparative Analysis:

#### - Time vs. Matrix Size:

\* GMRES shows a near-linear increase in computational time with the size of the matrix. This performance is remarkable, considering that iterative methods can be sensitive to matrix conditioning. The near-linear relationship might be a consequence of effective preconditioning or the specific matrix structures enabling more efficient computation.

#### - Iterations vs. Matrix Size:

\* The number of iterations needed by GMRES also increases linearly with matrix size. This is typical for iterative methods, where the number of iterations needed tends to increase with problem size but is also influenced by the conditioning of the matrix and the effectiveness of any preconditioner used.

#### - Accuracy vs. Matrix Size:

\* The accuracy graph suggests that the solutions become less accurate as the matrix size increases. This is an expected trend for high condition number matrices, as the error is more likely to propagate and accumulate over iterations in large-scale problems.

#### - Conclusition:

- \* For high condition numbers, both algorithms scale similarly in time with increasing matrix size, with GMRES showing a slight advantage in iterations.
- \* The decrease in accuracy with matrix size is a clear indication that both algorithms are affected by the high condition number, though this effect seems to be slightly less severe for GMRES.
- \* The results suggest that while GMRES maintains its stability in terms of iterations required, LU-IR may have limitations in the accuracy and effectiveness of its iterative refinements as the matrix size grows.

## References

- [1] Azzam Haidar, Harun Bayraktar, S.T.J.D., Higham, N.J.: Mixed-precision iterative refinement using tensor cores on gpus to accelerate solution of linear systems. Proc.R.Soc.A 476:20200110 (2020)
- [2] Boris I. Krasnopolsky, A.V.M.: Evaluating pereformance of mixed precision linear solvers with iterative refinement. Supercomputing Frontier and Innovations (2021)
- [3] F.Walker, H.: Implementation of the gmres method using householder transformations. SIAM J. Sci. STAT.Comput. (January 1988)
- [4] Jennifer A. Loe, Christian A. Glusa, I.Y.E.G.B.S.R.: Experimental evaluation of multiprecision strategies for gmres on gpus. IEEE IPDPS Accelerators and Hybrid Emerging Systems (AsHES) (2021)