1 PRECONDITIONING VIA GMRES IN POLYNOMIAL SPACE *

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Abstract. We propose a class of polynomial preconditioners for solving non-Hermitian linear 3 4 systems obtained from a least-squares approximation in polynomial space instead of a standard Krylov subspace. The process for building the polynomial relies on an Arnoldi-like procedure in a 5 6 small dimensional polynomial space and is equivalent to performing GMRES in polynomial space. It is inexpensive and produces results with superior numerical stability. A few improvements to 8 the basic scheme are discussed including the development of a short-term recurrence and the use of compounded preconditioners. Numerical experiments, including a test with challenging 3D Helmholtz 9 10 equations and a few publicly available sparse matrices, are provided to demonstrate the performance 11 of the proposed preconditioners.

12 **Key words.** Polynomial preconditioning, polynomial iteration, orthogonal polynomial, short-13 term recurrence, Helmholtz equation

14 **AMS subject classifications.** 15A06, 49M25, 65F08, 65F10, 65F50

15 **1. Introduction.** We consider solving a large non-Hermitian linear system of 16 equations

17 (1.1)
$$Ax = b,$$

where $A \in \mathbb{C}^{N \times N}$ is non-Hermitian and $x, b \in \mathbb{C}^N$. A Krylov subspace method accel-18 erated by a certain type of preconditioner is often preferred for this type of problems, 19 e.g., GMRES with a form of the incomplete LU (ILU) factorization. However, when 20 the coefficient matrix A is highly indefinite (eigenvalues of A appear on both sides of 21 the imaginary axis) or extremely ill-conditioned, this method may suffer from slow 22convergence or even stagnation due to stability issues [25]. Furthermore, since both 23the construction and application phases of ILU preconditioners are sequential in na-24ture and lack of parallelism, ILU preconditioners cannot easily take full advantages 25of modern high-performance computing architectures such as distributed memory 26machines or GPUs. Recently, a class of preconditioners based on low-rank approxi-27mations has been developed to overcome these difficulties [17, 19, 34]. These precon-28ditioners explore the recursive or hierarchical low-rank approximation of the Schur 29 complement or its inverse and only apply ILU to the diagonal blocks in the reordered 30 matrix. In particular, the generalized multilevel Schur complement low-rank (GM-31 SLR) preconditioner [9] has been shown to be quite effective for both non-symmetric 32 and indefinite problems. 33

This paper discusses a new class of polynomial preconditioning techniques for solving (1.1). These preconditioners can be either used in a standalone way or they can be combined with those low-rank approximation type preconditioners to further improve efficiency.

Most classical acceleration schemes are in fact in the form of a polynomial iteration. Indeed, given an initial guess x_0 and residual $r_0 = b - Ax_0$, the approximate

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40 solution \tilde{x} in a given iteration is of the form:

$$\tilde{x} = x_0 + p(A)r_0,$$

42 where p is a polynomial, and its related residual is equal to

43 (1.2)
$$\tilde{r} = b - A\tilde{x} = (I - Ap(A))r_0 \equiv \rho(A)r_0$$

Note that the approximate solution is a member of the affine Krylov subspace x_0 + 44 $\mathcal{K}_m(A, r_0)$. The acceleration procedures based on Krylov subspace methods that 45have been developed in the literature are all based on polynomial iterations where 46 the iterates are of the form given above, and the polynomials are obtained using 47 48 various criteria. For example, the criterion employed in GMRES [26] is to select the polynomial p to make the residual norm $\|\tilde{r}\|_2$ as small as possible. The Chebyshev 49"semi-iterative" method [12, 13] constructs p so that the residual polynomial $\rho(t)$ is an 50 appropriately shifted and scaled Chebyshev polynomial of the first kind. The residual polynomial is built so that it is small in an ellipse that encloses the spectrum of the matrix A. In these methods, the polynomial p can be either used directly to solve 53 linear systems approximately in an iterative scheme as in [12, 13] and other works, or it can be exploited as a preconditioner in combination with an acceleration such as GMRES for example. 56

Polynomial preconditioners are quite appealing because they are simple to use 57and because they can be highly effective for some problems. The construction of the 58 polynomial preconditioner does not involve matrix factorizations and it is also independent of reordering schemes. Moreover, applying the preconditioner relies heavily 60 on one single operation namely the matrix-vector multiplication associated with the 61 original coefficient matrix A. This operation has been studied and optimized for over 62 decades by researchers, see, e.g., [2, 32, 3, 18] and is often extremely efficient for 63 sparse matrices. In addition, the computations are completely free of inner product 64 65 which is communication-intensive and limits the performance in a distributed memory environment. The paper brings three main contributions which are summarized 66 below: 67

- Improved numerical stability. In the past, several polynomial precondi-68 tioners have been proposed in the literature [24, 22, 21, 11]. However, all of 69 these methods suffer from numerical stability issues that hampers their use 70 71 for higher degrees. In contrast, the proposed methods build a polynomial basis via an Arnoldi-like procedure. This procedure represents the polynomial 72 implicitly and has well-controlled numerical stability. As a result, the pro-73 posed polynomial preconditioners can be computed accurately for arbitrary 7475 degree.
 - Guaranteed effect in spectrum. The proposed polynomial preconditioners are constructed by solving a discrete least-squares problem based on the spectrum of the coefficient matrix so that the spectrum of the preconditioned system will be better clustered. In contrast, those based on GMRES polynomials cannot guarantee to yield a good preconditioner as pointed out in [31, 11].
- Efficient construction and application. The proposed polynomial preconditioners are built in a carefully designed polynomial space which has much smaller dimension compared to the matrix size. As a result, the cost of building the polynomial is essentially negligible. In the application phase, a technique based on short-term recurrence is proposed in Section 3.3 which

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can significantly accelerate the application of the preconditioner on a vectorand reduce the storage requirement.

The rest of this paper is organized as follows. Section 2 introduces a few ways to derive polynomial preconditioners based on solving minimax problems. Section 3 presents an Arnoldi-like procedure to generate a stable polynomial basis based on the boundary of the spectrum of the coefficient matrix. Several improvements are discussed in Section 4 and numerical examples are provided in Section 5. Finally, concluding remarks are draw in Section 6.

95 **2.** Polynomial construction via an explicit basis. In this section, we will 96 discuss a few ways to derive a polynomial preconditioner when an explicit basis $\{\phi_i(z)\}$ 97 for the polynomial space is given.

2.1. Classical minimax problem. In many applications, the boundary of the spectrum of A is not hard to estimate. For example, this can be done either by analyzing the physical problem [20] where (1.1) is derived from or approximated by methods such as the Arnoldi iteration [1, 10]. Assume that all eigenvalues of A are contained in a simply connected domain $\Omega \subset \mathbb{C}$ and denote by $\Gamma = \partial \Omega$ the boundary of Ω . Here, we further assume that Ω does not contain the origin and that Γ is piecewise smooth; see Figure 2.1 for an illustration.



FIG. 2.1. Eigenvalues of the matrix enclosed by a closed curve.

105 From (1.2) we have that

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$$\|\tilde{r}\| \le \|I - Ap(A)\| \|r_0\|$$

In order to make $\|\tilde{r}\|$ small, we could choose p so that $\|I - Ap(A)\|$ is small. A 107108 straightforward criterion to ensure this is simply to require that |1 - zp(z)| be small for all $z = \lambda$ where λ is an eigenvalue of A. Unfortunately, this approach involves 109110 all the eigenvalues of A, which is not practically feasible so an alternative is to seek p so that the maximum of |1 - zp(z)| in the region Ω is small. Since we assume the 111 eigenvalues of A are enclosed by Γ , and since 1 - zp(z) is holomorphic, the maximum 112 modulus principle [27] tells us that the maximum value of |1-zp(z)| on Ω is achieved 113 114 on the boundary Γ . Thus for a fixed m > 0, the sought-after polynomial p can be 115 characterized by the following minimax problem:

116 (2.1)
$$\min_{p \in \mathcal{P}_{m-1}} \max_{z \in \Gamma} |1 - zp(z)|,$$

117 where \mathcal{P}_{m-1} denotes the set of all complex polynomials of degree less than m.

It is important to note that an approach based on this framework can be viewed as a heuristic only because in the highly non-normal case the norm of ||I - Ap(A)|| is not always tightly related to the maximum of |1 - zp(z)| on the contour Γ that contains the spectrum, see, for example, the articles on the Crouzeix conjecture [6, 7, 8]. For many practical problems minimizing some norm of |1 - zp(z)| on the contour Γ will yield good results.

124 Defining the Chebyshev norm on any set $\mathcal{D} \subset \mathbb{C}$ of a function f by $||f||_{\mathcal{D}} =$ 125 $\max_{z \in \mathcal{D}} |f(z)|$, the minimax problem (2.1) can be rewritten as

126 (2.2)
$$\min_{p \in \mathcal{P}_{m-1}} \|1 - zp(z)\|_{\Gamma}.$$

127 This is a Chebyshev approximation problem in functional form with a domain that is 128 a continuous subset of the complex plane. The problem can be solved by a Remez-like 129 algorithm [4, 30, 23] or the Lanczos τ -method [15, 5]. However, when the geometry of 130 Γ becomes irregular or the degree of the polynomial increases, these methods might 131 fail. As a result, we will not attempt to solve the minimax problem (2.1) directly.

We can instead solve a discrete version of the problem, i.e., we can simplify (2.1)132by replacing the continuous contour Γ by a discrete one. Let $\Gamma_n = \{z_1, z_2, \ldots, z_n\}$ 133 be an *n*-point discretization of the boundary Γ . This discretization should capture 134 the geometric characteristics of Γ , a uniform discretization of Γ usually suffices in 135practice. In certain cases when Γ contains a high curvature or discontinuous part, 136we can either add additional points to refine the discretization in this area or simply 137replace this part by a smoother curve before the discretization. We then consider 138 the Chebyshev norm on the discrete set Γ_n and define the following discrete minimax 139problem: 140

141 (2.3)
$$\min_{p \in \mathcal{P}_{m-1}} \|1 - zp(z)\|_{\Gamma_n}.$$

Write $p(z) = \sum_{i=1}^{m} \alpha_i \phi_i(z)$ and denoted by $\alpha = [\alpha_1, \alpha_2, \dots, \alpha_k]^T \in \mathbb{C}^m$ the column vector of all the coefficients, (2.3) becomes

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$$\min_{\alpha \in \mathbb{C}^m} \max_{1 \le i \le n} \left| 1 - z_i \sum_{j=1}^m \alpha_j \phi_j(z_i) \right|.$$

145 Define an $n \times m$ matrix F with entries given by

146
$$f_{ij} = z_i \phi_j(z_i), \quad 1 \le i \le n, \ 1 \le j \le m,$$

147 and $e \in \mathbb{C}^n$ the column vector of all ones, (2.3) can be reformulated in the matrix 148 form as

149
$$\min_{\alpha \in \mathbb{C}^m} \|e - F\alpha\|_{\infty}.$$

We refer the readers to [29, 28, 33, 16] for some discussions on algorithms for solving the above complex linear programming problem. This problem uses the infinity norm in \mathbb{C}^n . We will not consider this approach in the remainder of the paper.

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Instead we will replace the infinity norm by the 2-norm in \mathbb{C}^n . The least-squares polynomial will be computed by a GMRES-like procedure in polynomial space which is described next.

3. Polynomial construction via an Arnoldi process. Define an inner prod uct for the polynomial space as

158 (3.1)
$$\langle p_1, p_2 \rangle = \sum_{i=1}^n p_1(z_i) \overline{p_2(z_i)}.$$

This sesqui-linear form is a valid inner product of the space of polynomials \mathcal{P}_m as long as *m* does not exceed the number of points *n*. We will denote by $\|\cdot\|_{\omega}$ the related norm. Then we would like to solve the following discrete least-squares problem instead of (2.2):

163 (3.2)
$$\min_{p \in \mathcal{P}_{m-1}} \|1 - zp(z)\|_{\omega}^2.$$

Instead of specifying a basis $\{\phi_i(z)\}_{i=1}^m$ in advance as in Section 2, we will actually build the polynomial basis dynamically in an Arnoldi-like a process.

3.1. GMRES in polynomial space. The construction procedure for the optimal polynomial is similar to GMRES in vector space and is described in Algorithm 3.1. For the sake of conformity with the notation used in the standard Arnoldi process, the polynomial basis of degree l is represented by q_{l+1} , instead of q_l .

It is easy to see that the Arnoldi-like process Algorithm 3.1 will indeed generate 170 a set of orthonormal polynomial basis $\{q_i\}_{i=1}^m$ with respect to the inner product (3.1), 171there will be no stability issue even for high degrees due to the full orthogonalization. 172The question now is how to represent the polynomials and how to carry out the actual 173computations that are involved in Algorithm 3.1. In fact we have a number of choices 174175of which we will only retain one. The simplest choice, a poor one for obvious reasons of stability, is to use the power series representation. In this case, a polynomial p(z) =176 $\alpha_0 + \alpha_1 z + \cdots + \alpha_{m-1} z^{m-1}$ will be represented by the vector $[\alpha_0, \alpha_1, \cdots, \alpha_{m-1}]^T \in \mathbb{C}^m$. 177For example, the polynomial multiplication $q := zq_i$ in Step 3 amounts to shifting 178all components of the representing vector down by one position and putting a zero 179in the first position; addition, subtraction and scalar multiplication all translate to 180the corresponding operation on the vector; inner products are also easy to compute 181 efficiently once the Gram matrix of the power series basis is computed. 182

However, we will not use any explicit representations because, as we will show 183 later, we are more interested in the coefficients h_{ii} than the polynomials themselves. 184Therefore we will represent the polynomials implicitly by the evaluations on the points 185 $\{z_i\}_{i=1}^n$, i.e., a polynomial p is represented by a vector $[p(z_1), p(z_2), \ldots, p(z_n)]^T \in \mathbb{C}^n$. 186 Under this representation, the polynomial multiplication $q := zq_i$ in Step 3 will be 187 translated simply into the entry-wise multiplication of two vectors of length n, and 188 the inner products in Steps 5 and 8 become standard inner products in vector space 189 $\mathbb{C}^{n}.$ 190

We now address the solution of the discrete least-squares problem (3.2). Define the $(m + 1) \times m$ matrix H_m where $(H_m)_{ij} = h_{ij}$, for $i \leq j + 1$ and $(H_m)_{ij} = 0$, for i > j + 1, so that H_m is an upper-Hessenberg matrix. If we abuse the notation and replace all polynomials by their vector representations in Algorithm 3.1, then the constant 1 in (3.2) becomes βq_1 where $\beta = ||\mathbf{1}||_{\omega} = \sqrt{n}$. Define $Q_l = [q_1, q_2, \dots, q_l]$ to be the column concatenation of the first l basis vectors, then each Q_l , for all

Algorithm 3.1 The Arnoldi-like process in polynomial space

Input: Discretization points $\{z_i\}_{i=1}^n$ on Γ and degree mOutput: Orthogonal polynomial basis $\{q_i\}_{i=1}^{m+1}$ 1: Set $q_1 = 1/||1||_{\omega}$ $\triangleright q_1$ is of degree 0 and norm 1 2: for j = 1, 2, ..., m do Compute $q := zq_j$ \triangleright Increase degree 3: for i = 1, 2, ..., j do 4: 5:Compute $h_{ij} = \langle q, q_i \rangle$ Compute $q = q - h_{ij}q_i$ 6: \triangleright Full orthogonalization 7: end for Compute $h_{j+1,j} = ||q||_{\omega}$ 8: Compute $q_{j+1} = q/h_{j+1,j}$ \triangleright Normalize the new basis 9: 10: end for

197 $1 \le l \le m+1$, is unitary. If p is expressed linearly in the basis $\{q_1, q_2, \ldots, q_m\}$ as

$$p = \sum_{i=1}^{m} \alpha_i q_i = Q_m \alpha$$

199 where $\alpha = [\alpha_1, \alpha_2, \dots, \alpha_m]^T$, then the polynomial zp in (3.2) becomes

$$zp = \sum_{j=1}^{m} \alpha_j (zq_j) = \sum_{j=1}^{m} \alpha_j \sum_{i=1}^{j+1} h_{ij} q_i = \sum_{j=1}^{m} \alpha_j \sum_{i=1}^{m+1} h_{ij} q_i \quad (h_{ij} = 0 \text{ when } i - j > 1)$$
$$= \sum_{i=1}^{m+1} \sum_{j=1}^{m} q_i h_{ij} \alpha_j = Q_{m+1} H_m \alpha.$$

In the end we observe that solving (3.2) amounts to minimizing with respect to $\alpha \in \mathbb{C}^n$ the objective function

203
$$J(\alpha) = \|\beta q_1 - Q_{m+1} H_m \alpha\|_2^2$$

Since Q_{m+1} is unitary and $q_1 = Q_{m+1}e_1$ where $e_1 = [1, 0, ..., 0]^T$ is a vector of length m + 1, this can be further reduced to

206 (3.3)
$$J(\alpha) = \|\beta e_1 - H_m \alpha\|_2^2.$$

Note that this is a standard least-squares problem, which is identical to the one encountered in the GMRES process.

209 Once α is found from (3.3), we obtain a polynomial p of degree m-1 and the 210 matrix M defined by $M^{-1} = p(A)$ can be used as a preconditioner for solving the 211 linear system Ax = b.

To apply M^{-1} to a vector v, note that

213 (3.4)
$$M^{-1}v = p(A)v = \sum_{i=1}^{m} \alpha_i q_i(A)v := \sum_{i=1}^{m} \alpha_i v_i$$

214 where we define $v_i \equiv q_i(A)v$ for $1 \leq i \leq m$. Since $q_1 = 1/\sqrt{n}$ so

215 (3.5)
$$v_1 = q_1(A)v = Iv/\sqrt{n} = v/\sqrt{n}.$$

From the Arnoldi-like process Algorithm 3.1 we have that $zq_i = \sum_{j=1}^{i+1} h_{ji}q_j$, thus

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$$Av_i = Aq_i(A)v = \left[\sum_{j=1}^{i+1} h_{ji}q_j(A)\right]v = h_{i+1,i}v_{i+1} + \sum_{j=1}^{i} h_{ji}v_j, \quad 1 \le i \le m-1.$$

and hence

219 (3.6)
$$v_{i+1} = \frac{1}{h_{i+1,i}} \left(Av_i - \sum_{j=1}^i h_{ji}v_j \right), \quad 1 \le i \le m-1.$$

The v_i 's can be computed recursively from (3.5) and (3.6) and the final result is just a linear combination of v_i 's with the coefficient α . Note that the only information needed is the pre-calculated entries available in H_m , the basis Q_{m+1} is not involved directly in the least-squares problem (3.3) for finding α or in applying the preconditioner (3.4)–(3.6).

225We note that the idea of using an Arnoldi-like procedure to generate orthogonal polynomials is not completely new. In fact, the framework is similar to what was 226discussed in [24] where the Chebyshev polynomial basis is used to construct a poly-227 nomial p that minimizes the residual polynomial 1 - zp under some specially defined 228norm. But, as mentioned in [24], this algorithm suffers from numerical stability issues 229 230 and the polynomial degree has to be kept low. The main reason is that the two norms 231 used in [24] are completely different, namely, the norm used to form the Chebyshev polynomial basis and the one used to characterize the solution are not compatible. As 232 a result, the orthogonal polynomial basis is no longer orthogonal in the inner product 233space associated with the optimization problem that generates the solution. The same 234235argument holds true for other methods that try to construct a polynomial from the 236 span of a given basis. Since the algorithm proposed in this manuscript uses only the inner product (3.1) and implicitly constructs the polynomial p, p will be accurately 237 computed for high degrees (as long as m < n). Another class of method construct the 238polynomial by finding all of its roots and represents the polynomial by the product 239 240of a series of degree one polynomials, e.g., in [21, 11]. These methods also suffer from stability issues when the degree is high mainly due to numerical cancellation. 241

242 **3.2.** Connection to GMRES. In comparing the proposed approach to the standard GMRES approach, one can observe that (3.3) is exactly the same least-243244squares problem that we solve in standard GMRES except that the coefficients of H_m are generated in a vector space of dimension n, the number of points on the contour. 245Looking more carefully at the algorithm, it is also possible to show that in fact *it* 246is equivalent to the standard GMRES algorithm applied to the diagonal matrix whose 247entries are the discretization points z_1, z_2, \dots, z_n . They are equivalent in the sense 248 that they would generate the same Hessenberg matrix H_m and in the end also the 249same polynomial. For this reason we may refer to this approach as a *proxy-GMRES* 250algorithm since that the original matrix is replaced by a small ("proxy") diagonal 251matrix whose spectrum captures the original spectrum well. 252

3.3. Short-term recurrence. Because the matrix H_m in Algorithm 3.1 is an upper Hessenberg matrix, computing p(A)v for a degree m-1 polynomial p costs $\mathcal{O}(m^2N)$ operations and requires $\mathcal{O}(mN)$ storage. This implies that despite the good numerical stability of the algorithm, its computation cost and storage quickly become unacceptably high as m increases. Motivated by the three-term recurrence for Cheby-

shev polynomials, we will show in this section that a short-term recurrence can be exploited to significantly reduce these costs.

The basic idea is to replace the full orthogonalization in Steps 4 to 7 in Algorithm 3.1 by a partial orthogonalization. That is, the newly generated polynomial qin Step 3 is only orthogonalized against the most recent k basis, which leads to the following short-term recurrence relation

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$$t_{j+1,j}\hat{q}_{j+1} = z\hat{q}_j - \sum_{i=j-k+1}^j t_{ij}\hat{q}_i, \quad 1 \le j \le m,$$

where t_{ij} $(1 \le i \le j)$, $t_{j+1,j}$ and \hat{q}_{j+1} are generated in the same way as in Steps 5, 8 and 9 in Algorithm 3.1, respectively. The computed basis $\{\hat{q}_i\}_{i=1}^{m+1}$ form the columns of \hat{Q}_{m+1} and t_{ij} 's form an $(m+1) \times m$ matrix T_m . Notice that \hat{Q}_{m+1} is not unitary anymore and T_m is a banded matrix with one subdiagonal and k-1 superdiagonals. For example, when k = 2, we have the three-term recurrence for the computed basis \hat{q}_i and T_m is a tridiagonal matrix. This is similar to the Chebyshev polynomial case. In the extreme case when k = m, the partial reorthogonalization becomes equivalent to the full orthogonalization and all results in Section 3.1 are recovered.

to the full orthogonalization and all results in Section 3.1 are recovered. Similar to Section 3.1, with the new basis $\{\hat{q}_j\}_{j=1}^{m+1}$ from the short-term recurrence we can rewrite (3.2) into minimizing with respect to $\hat{\alpha} \in \mathbb{C}^n$ a new objective function

275 (3.7)
$$\hat{J}(\hat{\alpha}) = \|\beta \hat{q}_1 - \hat{Q}_{m+1} T_m \hat{\alpha}\|_2^2.$$

Solving for $\hat{\alpha}$ in this problem typically needs to compute an orthogonal factorization of the matrix $\hat{Q}_{m+1}T_m$, which requires some additional computation cost and storage (since both \hat{Q}_{m+1} and T_m need to be stored) compared to computing α in (3.3). However, recall that all these computations are still within a vector space of dimension n which is typically much smaller then N.

On the other hand, applying the preconditioner $M^{-1} = \hat{p}(A)$ where \hat{p} is represented in the new basis \hat{Q}_m is slightly different. More specifically, (3.6) is replaced by the corresponding short-term version

284 (3.8)
$$v_{i+1} = \frac{1}{t_{i+1,i}} \left(Av_i - \sum_{j=i-k+1}^i t_{ji} v_j \right).$$

Due to the above short-term recurrence, the application of the preconditioner $M^{-1} = \hat{p}(A)$ on a vector only requires $\mathcal{O}(mkN)$ operations and $\mathcal{O}(kN)$ storage.

Now we discuss the stability issue associated with this approach. In exact arithmetics, it is easy to see that (3.3) and (3.7) are equivalent and the polynomials $p = Q_m \alpha$ and $\hat{p} = \hat{Q}_m \hat{\alpha}$ obtained from both orthogonalization schemes are exactly the same. This is because enforcing a short-term recurrence is equivalent to a change of basis and an update to the corresponding coefficients. However, in floating point arithmetics, \hat{Q}_m becomes increasingly ill-conditioned when *m* increases and thus the coefficient $\hat{\alpha}$ becomes increasingly hard to compute accurately.

Next we study the relation between the conditioning of the basis matrix \hat{Q}_m and the number of recurrence terms k. Figure 3.1 shows how the 2-norm condition number $\kappa_2(\hat{Q}_m)$ grows for multiple values of k where Γ and Γ_n are draw from the numerical example in Section 5.2. In Figure 3.1a, the condition number plots for k = 2 and 3 almost coincide, the relative error between them is shown in Figure 3.1b; similarly for k = 4, 5 and k = 6, 7. When the recurrence is too short, i.e., k = 2 or 3, the condition number rapidly grows beyond 10^{12} when m passes 60. In that case, the polynomial \hat{p} solved with this basis becomes inaccurate and the resulting preconditioner may become useless. By increasing k to 4 or 5, $\kappa_2(\hat{Q}_m)$ quickly drops from 10^{12} to about 10^3 at m = 60 and the basis becomes too ill-conditioned again only when m reaches 180. Figure 3.1a also shows that the numerical stability keeps getting improved when k increases to 6.



FIG. 3.1. Conditioning of \hat{Q}_m generated with k-term recurrence.

The numerical stability of \hat{Q}_m can be monitored inexpensively by its associated Gram matrix. Denote by \hat{G}_m the $m \times m$ Gram matrix of the basis \hat{Q}_m whose entries are defined by

$$\hat{g}_{ij} = \langle \hat{q}_i, \hat{q}_j \rangle, \quad 1 \le i, j \le m$$

where the inner product is as defined in (3.1). The matrix \hat{G}_m is Hermitian positive definite. Let $\hat{G}_m = \hat{L}_m \hat{L}_m^H$ be the Cholesky factorization where \hat{L}_m is lower-triangular 310 311 and note that $\kappa_2(\hat{Q}_m) = \sqrt{\kappa_2(\hat{G}_m)} = \kappa_2(\hat{L}_m)$. As the Arnoldi-like process proceeds, 312 both the Gram matrix \hat{G}_{m+1} and the Cholesky factor \hat{L}_{m+1} can be quickly updated 313 with \hat{G}_m and \hat{L}_m from the previous step. When a high degree polynomial needs 314 to be used, the ill-conditioning of \hat{Q}_m can be quickly detected by keeping track of 315the condition number of the Cholesky factor \hat{L}_m . Whenever $\kappa_2(\hat{L}_m)$ goes beyond 316a certain tolerance, we can stop the process and accept the resulting polynomial 317 obtained at that point or restart the same process with a longer recurrence relation. 318 As is indicated in Figure 3.1, we can start from k = 2 and increase k by 2 every time 319 the process restarts. This process is repeated until the desired degree can be reached 320 while $\kappa_2(\hat{L}_m)$ still remains below the given tolerance. In practice, we find that setting 321 the tolerance at $\tau = 10^{12}$ usually yields good quality results. 322

4. Some improvements based on compounding preconditioners. In the previous sections, we always assume Γ will exclude the origin. This is because otherwise, the maximum modulus of the residual polynomial 1 - zp(z) will be no less than 1 on Γ and the resulting polynomial preconditioner will not be effective at all. For ill-conditioned problems, a few eigenvalues of A will stay in a small neighborhood of the origin and Γ has to be very close to the origin. In that case, a very high degree

polynomial becomes mandatory in order to keep the maximum value of |1 - zp(z)|on Γ strictly less than 1. On the other hand, the increased degree will require a longer recurrence and thus harm the efficiency of the proposed preconditioner. In this section, we will discuss two compounding techniques to overcome this difficulty.

4.1. Compounding two polynomials. The first approach is based on compounding two low degree polynomials to mimic the effect of a high degree polynomial. By doing this, even though the total number of matrix-vector multiplications associated with A might be slightly increased, the costs associated with vector operations and storage can be significantly reduced. Also as pointed out in [11], this strategy can also reduce the number of inner products performed.

This can be understood from a simple example. Suppose one high degree polyno-339 mial has degree m-1 and the other two low-degree polynomials have degree m_1-1 340 and $m_2 - 1$, respectively, with $m = m_1 \times m_2$. For these three polynomials, a k_i -term 341 recurrence is deployed for a polynomial of degree $m_i - 1$ (for i = 1, 2) while a k-term 342 recurrence is needed for degree m-1. Since m_1 and m_2 are both much smaller 343 344 than m, we have $k_1, k_2 \ll k$ when ensuring the numerical stability of the computed polynomial basis. Thus, applying the preconditioner resulting from compounding the 345 polynomials will entail fewer vector operations and storage. 346

We now provide more details on how to construct these two low-degree polynomi-347 als. First find a contour Γ that encloses all the eigenvalues of A and discretize it as Γ_{n_1} . 348 Based on Γ_{n_1} , construct the first polynomial p_1 of degree $m_1 - 1$ and select the recur-349 rence length k_1 with the procedure discussed in Section 3.3. It can be expected that 350 most of the eigenvalues of the preconditioned matrix $A_1 := M_1^{-1}A = p_1(A)A$ would 351be clustered around z = 1. Therefore, a second contour is then selected as a circle C352 centered at z = 1 with radius $\theta \in (0, 1)$. Let \mathcal{C}_{n_2} be an n_2 -point discretization of \mathcal{C} and 353 apply \mathcal{C}_{n_2} to compute the second polynomial p_2 with degree $m_2 - 1$ and recurrence 354length k_2 . In the end, the compound polynomial has the form $p(z) := p_1(z)p_2(zp_1(z))$ and the resulting preconditioner is $M^{-1} := p(A) = p_1(A)p_2(Ap_1(A))$. 355 356

It is clear that the preconditioner M^{-1} is a polynomial in A of degree $m_1m_2 - 1$. Applying M^{-1} on a vector involves two main operations:

- 1. Apply $p_1(A)$ to a vector, which follows the formula (3.4), (3.5), and (3.8). This computation costs $m_1 - 1$ matvecs associated with A and $\mathcal{O}(m_1k_1N)$ from vector operations and $\mathcal{O}(k_1N)$ storage;
- 362 2. Apply $p_2(Ap_1(A))$ to a vector. This operation consists of $m_2 1$ matvecs 363 of $Ap_1(A)$, $\mathcal{O}(m_2k_2N)$ extra costs from vector operations and $\mathcal{O}(k_2N)$ extra 364 storage.

Table 4.1 compares the costs of applying one high degree polynomial preconditioner verse one compound polynomial preconditioner. It is easy to see that when $m = m_1 \times m_2$, even though both preconditioners perform the same number of *matvecs* associated with A, the operations and peak storage associated with the compound polynomial preconditioner can be much less due to the fact that $k_1, k_2 \ll k$.

4.2. Compounding with other preconditioners. A second approach is to compound the polynomial preconditioner with other types of preconditioners. For ill-conditioned problems, it is suggested to perform an approximate factorization on $A + \sigma I$ for some complex shift σ [35] instead of the original coefficient matrix A. To simplify the discussion, we assume the incomplete LU factorization (ILU) is explored here:

TABLE 4.1

The cost and storage of applying the single and compound polynomial preconditioners, the single polynomial is of degree m-1, the compound polynomial is built with two low degree polynomials of degree $m_1 - 1$ and $m_2 - 1$.

	single polynomial	compound polynomial
matvec of A	m-1	$m_1 m_2 - 1$
vector operations	$\mathcal{O}(mkN)$	$\mathcal{O}(m_1m_2k_1N)$
peak storage	$\mathcal{O}(kN)$	$\mathcal{O}((k_1+k_2)N)$

$A + \sigma I \approx M_1 = LU.$ 376

377

We will now discuss two different ways to introduce a second level preconditioner. The first option is to compound M_1^{-1} with a polynomial preconditioner of the form p(A). Define the distance matrix E as $E = I - M_1^{-1}Ap(A)$. An ideal polynomial preconditioner of the distance matrix E as $E = I - M_1^{-1}Ap(A)$. 378 mial p should minimize $||E||_2$. Although its optimal solution is hard to calculate, an 380 approximate solution can be computed inexpensively by randomized sampling (see 381 [14] for details) as follows: first construct a set of polynomial basis $\{q_1, q_2, \ldots, q_m\}$ and express the polynomial as a linear combination of the basis $p = \sum_{j=1}^{m} \alpha_j q_j$, 382 383 then pick l random vectors $\omega_1, \omega_2, \ldots, \omega_l$ of length N and solve the coefficients $\alpha =$ 384 $[\alpha_1, \alpha_2, \ldots, \alpha_m]^T \in \mathbb{C}^m$ from the following problem 385

$$\min_{\alpha \in \mathbb{C}^m} \max_{1 \le j \le l} \|E\omega_j\|_2.$$

The main drawback of this approach is still numerical stability since it requires a pre-387 specified polynomial basis. No suitable norm like (3.1) can be defined in this case, 388 389 thus there is no reliable way to generate a good basis set like in Section 3.1.

390 The second option resolves this issue by considering a new linear system

391
$$M_1^{-1}Ax = M_1^{-1}b$$

and applying the procedures discussed in Section 3.1 to the new coefficient matrix 392 $A_1 := M_1^{-1}A$. Note here that the contour Γ for A_1 can be estimated by running a 393 few steps of the Arnoldi process. After the polynomial p is constructed, the com-394 pound preconditioner takes the form of $M^{-1} = p(M_1^{-1}A)$. Suppose the polynomial p is of degree m - 1, then one application of M^{-1} on a vector consists of m - 1395 396 matvecs associated with A and m-1 applications of the preconditioner M_1^{-1} . As 397 mentioned in Section 1, ILU type preconditioners may become the performance bot-398 tleneck in a parallel environment due to the sequential nature of the triangular solves. 399 This framework naturally allows replacing the ILU factorization with more scalable 400 preconditioners such as SLR, MSLR or GMSLR preconditioners [19, 34, 9]. 401

5. Numerical experiments. All numerical tests were ran in Matlab on a Desk-402 403 top PC with 3.80 GHz CPU and 8 GB memory. We used flexible GMRES (FGMRES) with a restart dimension of 50 as the accelerator, the initial guess was set to be the 404 405 zero vector and the process was terminated when either the residual was reduced by a prescribed factor τ or the total number of inner iterations reached 1000. 406

The following notation is used in this section: 407

• p-t: the computation time in seconds for constructing the preconditioner, 408which includes the time for estimating/discretizing the contour Γ , building 409

410	the polynomial or/and other preconditioners depending on the specific tests.
411	F indicates the preconditioner cannot be constructed;
412	• i-t: iteration time in seconds for FGMRES(50) to converge;
413	• its: total number of iterations required for FGMRES(50) to converge, F in-
414	dicates FGMRES(50) does not converge within 1000 inner iterations;

• mv: number of matvecs associated with A performed.

5.1. A diagonal matrix. In the first example, we generated a 2000×2000 416 diagonal matrix where all the diagonal entries (eigenvalues) were randomly chosen 417 from the semiannular region $\Omega = \{z \in \mathbb{C} \mid 0.8 \le |z| \le 2, 0 \le \operatorname{Arg}(z) \le \pi\}$. The 418 boundary of this region is shown in Figure 5.1 where the squares are the approximate 419 eigenvalues (Ritz values) computed by running 60 steps of the Arnoldi algorithm. An 420 421 approximate boundary was obtained by running Matlab's built-in function boundary on the approximate eigenvalues. Figure 5.1 shows that the Ritz values from the 422 Arnoldi algorithm can characterize the boundary of the spectrum. 423



FIG. 5.1. The exact and approximate boundaries of the spectrum and the approximate eigenvalues obtained from 60 steps of the Arnoldi algorithm for the $2,000 \times 2,000$ diagonal matrix in Section 5.1.

We first constructed polynomials of degree 29 (m = 30) with a recurrence length 424 425 k = 2. Figure 5.2 shows the contour maps for the function |1 - zp(z)| in log scale based on both exact (left) and approximate (right) boundaries. Since the estimated 426 boundary approximates the exact one very well, the two maps look almost identical. 427 Table 5.1 tabulates the numerical results for solving the linear system with these 428 constructed polynomial preconditioners, the tolerance was set at $\tau = 10^{-12}$. It took 429 237 iterations for FGMRES(50) to converge without any preconditioner. On the other 430 hand, FGMRES(50) with the polynomial preconditioners converged in 8 iterations in 431 432 both cases. Although the preconditioned methods performed 3 more matvecs, they actually took much less time to converge. Similar observations can also be made in 433 other examples in this section. This is due to the fact that a reduced iteration number 434 leads to a much smaller subspace for FGMRES and far fewer inner products during 435436 the computation. This performance gap can be expected to become more pronounced



437 when running the experiments on high performance computing architectures.

FIG. 5.2. Contour maps of |1-zp(z)| in log scale with different choice of Γ for the 2,000×2,000 diagonal matrix in Section 5.1. The asterisk marks the origin.

438 We also want to emphasize that the preconditioner construction time was only a

439 tiny fraction of the iteration time. This is because we used 400 discretization points

440 for both the exact and approximate boundaries and the corresponding polynomial

441 space has much smaller dimension compared to the matrix size N = 2000.

TABLE 5.1 Convergence results of FMGRES(50) for the 2,000 × 2,000 diagonal matrix test in Section 5.1 with tolerance $\tau = 10^{-12}$.

	p-t	i-t	its	mv	
no j	\	0.81	237	237	
with precord	exact boundary	0.0062	0.61	8	240
with precond.	approx. boundary	0.0056	0.61	8	240

442 **5.2. Helmholtz problem.** The second example is the 3D Helmholtz equation

$$-\Delta u - \frac{\omega^2}{c^2(x)}u = s$$

where ω is the angular frequency and c(x) is the wavespeed. The computational do-444 main was the unit cube and the equation was discretized with 7-point stencil finite 445 difference method. PML boundary conditions were imposed to reduce the artificial re-446 flections near the boundaries of the computational domain. We kept 8 grid points per 447 wavelength. The resulting linear system is sparse complex symmetric with dimension 448 $N = N_x \times N_y \times N_z$. Moreover, the spectrum of the matrix is contained in a rectangle 449area $\{z \in \mathbb{C} \mid \text{real}(z) \in [-1, \rho_1 - 1], \text{imag}(z) \in [-\rho_2, 0]\}$ where the two parameters ρ_1 450and ρ_2 are given by [20, Lemma 3.1]. Figure 5.3a shows all the eigenvalues and the 451rectangular boundary from [20, Lemma 3.1] for a discretized Helmholtz operator of 452 size $N = 20^3$. We fixed the tolerance at $\tau = 10^{-3}$ for the Helmholtz equation tests in 453this section. Note that this test matrix is non-normal. 454

455 **5.2.1. Compounding polynomial preconditioners.** Compared with the first 456 test example, this problem is much harder to solve. First, there are many eigenvalues



FIG. 5.3. The theoretical rectangular spectrum boundary from [20, Lemma 3.1] and the zoom-in view of the modified Γ near the origin for a discretized Helmholtz operator of size $N = 20^3$

close to the origin. Second, the theoretical spectrum boundary (the rectangular area) 457overlaps with the origin. In order to construct an effective polynomial preconditioner, 458we have to 459

460

462

1. Modify the theoretical rectangular contour [20, Lemma 3.1] to exclude the origin. 461

2. Use a high degree polynomial.

For this test, the problem size was $N = 100^3 = 1,000,000$ and the boundary 463 Γ was a modified rectangle with the origin excluded. A zoom-in view of Γ near the 464 origin is shown in Figure 5.3b. The boundary Γ was then discretized uniformly on 465each of its continuous parts with a step size h = 0.002 which results to a total of 466 3551 discretization points. We compared the performance of two polynomial precon-467 468 ditioners on this test matrix. The first one was a single polynomial of degree 599 (m = 600). In order to ensure its numerical stability, we used a recurrence length 469k = 10. The second one was a compound polynomial with $m_1 = 60$ and $m_2 = 10$ 470 so that $m_1 \times m_2 = m$. Since m_1 and m_2 are relatively small, the recurrence lengths 471were set to be $k_1 = k_2 = 2$. The convergence results with these two preconditioners 472are shown in Table 5.2. In addition, we also ran ILUT-preconditioned FGMRES(50) 473and CG on the corresponding normal equation with a block Jacobi preconditioner for 474comparisons. 475

Preconditioner type	p-t	i-t	its
no preconditioner		\	F
ILUT	F	\	
CG with diagonal preconditioner	0.49	\	F
single polynomial of degree $600 - 1$	5.85	2554.44	16
compound polynomial of degree $60 \times 10 - 1$	0.08	853.11	18

TABLE 5.2

Convergence results of various preconditioned FGMRES(50) on the Helmholtz equation test with size $N = 100^3$, the tolerance is fixed at $\tau = 10^{-3}$.

Due to the ill-conditioning and indefiniteness of the test matrix, the first three 476477 methods in Table 5.2 failed to converge. In particular, ILUT even failed to finish the factorization. On the other hand, both polynomial preconditioned methods converged within 18 iterations. Compared to the single polynomial preconditioner, the compound polynomial preconditioner took much less time to construct and reduced the iteration time by more than a half even though 1200 more matvecs of A were

482 performed.

The residual histories are plotted in Figure 5.4. It is easy to see that both polynomial preconditioned FGMRES methods converged quickly without encountering any stagnation while the non-preconditioned FGMRES converged slowly.



FIG. 5.4. Relative residual histories of three preconditioned FGMRES(50) on the Helmholtz equation test of size $N = 100^3$.

5.2.2. Compounding with SLR. We also compounded the polynomial pre-486 conditioner with the nonsymmetric SLR preconditioner [19] and tested its precondi-487 tioning effect on the Helmholtz problem. The problem size was still kept at $N = 100^3$ 488 and the tolerance was set at $\tau = 10^{-3}$. We applied the SLR preconditioner to the 489shifted system $M_1 \approx A + \sigma I$ with a complex shift $\sigma = -0.4i$ (pulling the eigenvalues 490away from the origin), and then chose a polynomial preconditioner of degree 29 for 491 the matrix $A_1 = M_1^{-1}A$. The approximate spectrum boundary of A_1 was obtained by 492 running 80 steps of Arnoldi process, then it was uniformly discretized with 730 points. 493 The convergence results as well as the comparison with SLR preconditioner are shown 494 495Table 5.3. We see that SLR preconditioned FGMRES(50) failed to converge in 1000 iterations while the SLR compound polynomial preconditioner converged in only 29 496 iterations and required the least iteration time among all seven methods tested in 497 Table 5.2 and Table 5.3. Also notice that the construction time of this compound 498 preconditioner is higher than other methods because it include both the SLR precon-499500ditioner construction time and the time to perform 80 steps of Arnoldi process.

In order to visualize the spectrum of the preconditioned matrix across different stages with this compound preconditioner, we also ran the experiment on a smaller Helmholtz problem of size $N = 20^3$ so that we were able to compute all eigenvalues of the matrices. Let M_1 denote the SLR preconditioner for the discretized Helmholtz operator A. The spectrum of $A_1 = M_1^{-1}A$ as well as the approximate eigenvalues from running 80 steps of the Arnoldi algorithm are shown in Figure 5.5a. A polyno-

TABLE 5.3

Convergence results of the SLR and SLR compound polynomial preconditioned FGMRES(50) on the Helmholtz equation test with size $N = 100^3$, the tolerance was fixed at $\tau = 10^{-3}$.

Preconditioner type	p-t	i-t	its
SLR preconditioner	86.94		F
SLR with polynomial of degree $30 - 1$	145.85	308.03	29

mial preconditioner $p(A_1)$ of degree 29 was constructed and the contour map of the corresponding residual polynomial |1-zp(z)| is drawn in Figure 5.5b. Compared with Figure 5.3b, it is easy to see that the SLR preconditioner pushed the eigenvalues of A_1 further away from the origin. Thus, a low-degree polynomial of p has already led to an efficient preconditioner, which is supported by both the contour map Figure 5.5b as well as the spectrum of the preconditioned matrix $A_1p(A_1)$ in Figure 5.5c.



(a) Exact and approximate eigenvalues of $M_1^{-1}A$ (b) Contour map of |1 - zp(z)| in log scale



(c) Eigenvalues of the final preconditioned matrix $A_1p(A_1)$

FIG. 5.5. Illustration of the preconditioning effect of both stages of the SLR compound preconditioner $p(M_1^{-1}A)$ on a small discretized Helmholtz equation test of size $N = 20^3$, where M_1 denotes the SLR preconditioner and p has degree 29.

	Inf	formation	of the	e test	sparse	mat	rices.	
name		Orde	r		nnz			_

Group/matrix name	Order	nnz	Origin
Rajat/rajat09	24,482	105,573	circuit simulation problem
Dehghani/light_in_tissue	29,282	406,084	electromagnetics problem
Goodwin/Goodwin_127	178,437	5,778,545	CFD problem
Kim/kim2	456,976	11, 330, 020	3D problem
Bourchtein/atmosmodd	1,270,432	8,814,880	CFD problem

TABLE 5.5

Convergence results of general sparse matrices with FGMRES(50) and tolerance $\tau = 10^{-10}$, all polynomials were of degree 39, column n shows the number of discretization points used on the spectrum boundary of A_1 .

matrix	ILUT			SLR compound polynomial			
mauna	p-t	i-t	its	n	p-t	i-t	its
rajat09	F	\		763	0.46	3.27	44
light_in_tissue	0.071	2.39	213	572	0.54	0.71	6
Goodwin_127	F	\	\	460	2.19	409.74	311
kim2	4.85	20.52	38	1493	18.72	7.69	2
atmosmodd	1.17	454.66	397	557	36.10	42.71	6

6. Conclusions. The primary distinction between the polynomial precondition-528 529 ing techniques introduced in this paper and existing techniques is their aim at controlling the numerical stability of the polynomial construction and the resulting iterative 530 process, while allowing the degree of the polynomial to be high. Using a high degree 531polynomial is very important in order to guarantee a good quality preconditioner that will produce convergence in a smaller number of outer iterations. The performance of 534 the method is significantly improved by a process which relies on a short-term recurrence. It is clear that a big appeal of the method is its potential for great performance 535 in a highly parallel, possibly GPU based, environment. The numerical experiments 536show that even in a scalar environment, the method can be effective in solving difficult 538 problems when other techniques fail.

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