

A block Chebyshev-Davidson method for linear response eigenvalue problems

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Abstract We present a Chebyshev-Davidson method to compute a few smallest positive eigenvalues and corresponding eigenvectors of linear response eigenvalue problems. The method is applicable to more general linear response eigenvalue problems where some purely imaginary eigenvalues may exist. For the Chebyshev filter, a tight upper bound is obtained by a computable bound estimator that is provably

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correct under a reasonable condition. When the condition fails, the estimated upper bound may not be a true one. To overcome that, we develop an adaptive strategy for updating the estimated upper bound to guarantee the effectiveness of our new Chebyshev-Davidson method. We also obtain an estimate of the rate of convergence for the Ritz values by our algorithm. Finally, we present numerical results to demonstrate the performance of the proposed Chebyshev-Davidson method.

Keywords Eigenvalue/eigenvector \cdot Chebyshev polynomial \cdot Davidson type method \cdot Convergence rate \cdot Linear response \cdot Upper bound estimator

Mathematics Subject Classifications (2010) 65F15 · 15A18

1 Introduction

In computational quantum chemistry and physics, the so-called *random phase approximation* (RPA) describes the excitation states (energies) of physical systems in the study of collective motion of many-particle systems [2, 25, 26]. It has important applications in silicon and other nanoscale materials, analysis of interstellar clouds [2, 3], polarizabilities [20], and finding the electronic excitation spectrum of a quantum many-fermion system [27]. One important question in RPA is to compute a few eigenpairs associated with the smallest *positive* eigenvalues of the following eigenvalue problem:

$$\mathscr{H}w := \begin{bmatrix} A & B \\ -B & -A \end{bmatrix} \begin{bmatrix} u \\ v \end{bmatrix} = \lambda \begin{bmatrix} u \\ v \end{bmatrix}, \tag{1.1}$$

where $A, B \in \mathbb{R}^{n \times n}$ are both symmetric matrices and $\begin{bmatrix} A & B \\ B & A \end{bmatrix}$ is positive definite. Through a similarity transformation, this eigenvalue problem can be equivalently transformed into [2, 3, 19]

$$Hz := \begin{bmatrix} 0 & K \\ M & 0 \end{bmatrix} \begin{bmatrix} y \\ x \end{bmatrix} = \lambda \begin{bmatrix} y \\ x \end{bmatrix}, \tag{1.2}$$

where K = A - B and M = A + B. The eigenvalue problem Eq. (1.2) was still referred to as the *linear response eigenvalue problem* (LREP) [2, 19, 27] and will be so in this paper, too.

The condition imposed upon A and B in Eq. (1.1) implies that both K and M are real symmetric and positive definite [2]. But there are cases where one of them may be indefinite [17]. Throughout this paper, we consider the more general case:

$$K$$
 and M are Hermitian, one of them is positive definite, and the other may be indefinite. (1.3)

For this general case, the interested quantities are the first few eigenvalues whose squares are the smallest few, together with their corresponding eigenvectors.



For simplicity, in the rest of this paper, we assume that M is positive definite and K is Hermitian but may be indefinite. Doing so loses no generality because otherwise we simply interchange the roles of K and M.

From Eq. (1.2), we have $Kx = \lambda y$ and $My = \lambda x$ and they together produce

$$KMy = \lambda^2 y, \tag{1.4a}$$

$$MKx = \lambda^2 x. \tag{1.4b}$$

The three eigenvalue problems: (1.2) of H, (1.4a) of KM, and (1.4b) of MK, are theoretically equivalent. That is, if any one of them is solved, the solutions to theother two can be constructed from the solved one with little effort [2, Theorem 2.1].

Researches on solving Eq. (1.2), or equivalently Eqs. (1.4a) and (1.4b), have been very active, see [3, section 1] and references therein. The goal of this paper is to extend the Chebyshev-Davidson method and its block version proposed in [30, 33] for symmetric/Hermitian eigenvalue problems. The new method is capable of tackling the more general case (1.3) that may have purely imaginary eigenvalues whereas previous methods as surveyed in [3, section 1] cannot.

The rest of this paper is organized as follows. In Section 2, we present our algorithm and the convergence rate estimate based on an upper bound estimator. The computable upper bound estimator for $\lambda_{\max}(KM)$ and the adaptive strategy are detailed in Section 3. Numerical results of our algorithm are presented in Section 4. Finally, some conclusions are drawn in Section 5.

Throughout this paper, $\mathbb{K}^{n \times m}$ is the set of all $n \times m$ matrices with entries in \mathbb{K} , where \mathbb{K} is \mathbb{C} (the set of complex numbers) or \mathbb{R} (the set of real numbers), $\mathbb{K}^n = \mathbb{K}^{n \times 1}$, and $\mathbb{K} = \mathbb{K}^1$. I_n (or simply I if its dimension is clear from the context) is the $n \times n$ identity matrix, and e_j is its jth column. MATLAB-like convention is adopted to access the entries of vectors and matrices. For a matrix X, $X_{(:,i:j)}$ denotes the submatrix consisting the i-th to the j-th columns of X, diag(X) denotes the column vector of the diagonal entries of X, and eig(X) denotes the spectrum of X. If X is nonsingular, $\kappa_2(X)$ denotes the spectral condition number of X. The superscript "·H" takes conjugate transpose while "·T" takes transpose only. The number $\bar{\alpha}$ is the conjugate of the scalar α .

2 Chebyshev filter for LREP

2.1 Chebyshev filter

Chebyshev polynomials play important roles in theoretical analysis of numerical algorithms as well as in algorithmic design [21]. They are also useful in practice, as a means of accelerating single vector iterations or projection processes. In [30, 33], the polynomials are used to magnify the components of the desired eigenvectors while suppress those of the undesired ones.

The *m*-th Chebyshev polynomial of the 1st kind is

$$\mathcal{T}_m(t) = \begin{cases} \cos(m \arccos(t)), & \text{for } -1 \le t \le 1, \\ \cosh(m \arccos(t)), & \text{for } |t| \ge 1. \end{cases}$$
 (2.1)



It frequently shows up in numerical analysis and computations because of its numerous nice properties; for example, $|\mathcal{T}_m(t)| \le 1$ for $|t| \le 1$ and $|\mathcal{T}_m(t)|$ grows extremely fast¹ for |t| > 1. Also important is the associated three-term recurrence relation: $\mathcal{T}_0(t) = 1$, $\mathcal{T}_1(t) = t$, and for $m \ge 1$

$$\mathscr{T}_{m+1}(t) = 2t\,\mathscr{T}_m(t) - \mathscr{T}_{m-1}(t),\tag{2.2}$$

which is responsible for the numerical efficiency in its numerous applications.

For an $n \times n$ Hermitian matrix A, suppose $\operatorname{eig}(A) \subset [\alpha_0, \beta]$ and we are interested in the eigenvalues lying in $[\alpha_0, \alpha]$, where $\alpha_0 < \alpha < \beta$. The basic idea of the Chebyshev filtering approach is to use Chebyshev polynomials to magnify the components of an approximate eigenvector u in the directions of the wanted eigenvectors and at the same time suppress those in the directions of the unwanted eigenvectors. To this end, we first transform $[\alpha, \beta]$ onto [-1, 1] by the following affine mapping:

$$t \in [\alpha, \beta] \quad \rightarrow \quad \phi(t) = \frac{2t - (\alpha + \beta)}{\beta - \alpha} \in [-1, 1].$$
 (2.3)

Then $|\mathcal{T}_m(\phi(t))| \leq 1$ for $t \in [\alpha, \beta]$ and $|\mathcal{T}_m(\phi(t))|$ grows extremely fast for $t \notin [\alpha, \beta]$. For this reason, $\mathcal{T}_m(\phi(A))u$ magnify the components of u in the directions of the eigenvectors associated with eigenvalues in $[\alpha_0, \alpha)$ and at the same time suppress those in the directions of the eigenvectors associated with eigenvalues in $[\alpha, \beta]$. The effect is more dramatic as m increases. The three-term recurrence relation (2.2) yields

$$\mathscr{T}_{m+1}(\phi(A))u = 2\phi(A) \cdot \mathscr{T}_m(\phi(A))u - \mathscr{T}_{m-1}(\phi(A))u, \tag{2.4}$$

making it very efficient to compute $\mathcal{T}_m(\phi(A))u$.

What we just described is the basis of the Chebyshev filter in [30] and [33] for the large scale symmetric/Hermitian eigenvalue problem. But it is not difficult to notice that the idea can be applied to diagonalize matrices with real spectra. The following lemma [2] establishes some theoretical results, which show that Eq. (1.2) may be transformed into eigenvalue problems with real spectra. Therefore we can achieve acceleration by constructing appropriate Chebyshev filters.

Lemma 2.1 The following statements hold true for any matrices M and K in $\mathbb{C}^{n \times n}$, where M is positive definite and K is Hermitian.

(a) There exists a nonsingular $Y = [y_1, y_2, ..., y_n] \in \mathbb{C}^{n \times n}$ such that

$$K = Y\Lambda^2 Y^{\mathrm{H}}, \quad M = XX^{\mathrm{H}}, \tag{2.5}$$

where $\Lambda = \operatorname{diag}(\lambda_1, \lambda_2, \dots, \lambda_n)$, $\lambda_1^2 \leq \lambda_2^2 \leq \dots \leq \lambda_n^2$ and $X = Y^{-H} = [x_1, x_2, \dots, x_n]$.

 $[x_1, x_2, \dots, x_n].$ (b) $[\lambda_i y_i^H, x_i^H]^H$ is an eigenvector corresponding to λ_i of the matrix H,

$$H\begin{bmatrix} \lambda_i y_i \\ x_i \end{bmatrix} = \lambda_i \begin{bmatrix} \lambda_i y_i \\ x_i \end{bmatrix}. \tag{2.6}$$

¹In fact, a result due to Chebyshev himself says that if p(t) is a polynomial of degree no greater than m and $|p(t)| \le 1$ for $-1 \le t \le 1$, then $|p(t)| \le |\mathcal{T}_m(t)|$ for any t outside [-1, 1] [6, p.65].



- (c) Let (λ_i, z_i) (i = 1, 2) be two eigenpairs of H, and partition $z_i = [y_i^H, x_i^H]^H$. Then,
 - (i) if $\lambda_1 \neq \bar{\lambda}_2$, then $y_1^H x_2 + y_2^H x_1 = 0$.
 - (ii) if $\lambda_1 \neq \pm \lambda_2$, then $y_1^H x_2 = y_2^H x_1 = 0$.
- (d) Both KM and MK are diagonalizable. In fact,

$$KM = Y\Lambda^2 Y^{-1}, \quad MK = X\Lambda^2 X^{-1}.$$

Here M is assumed to be positive definite. If in addition K is positive semi-definite or definite, then $\lambda_1^2 \ge 0$ and thus we can make all $\lambda_i \ge 0$. In this case the eigenvalues of H are all real $\pm \lambda_i$,

$$-\lambda_n \le \dots \le -\lambda_2 \le -\lambda_1 \le \lambda_1 \le \lambda_2 \le \dots \le \lambda_n. \tag{2.7}$$

When K is indefinite, the first few λ_i^2 may be negative. To define λ_i unique, we take λ_i to be the unique square root of λ_i^2 with nonnegative imaginary part. Lemma 2.1 was proved for K being positive definite [2], but the proof carries over for indefinite K with only minor changes.

Often in LREP (1.2), both K and M are definite and the first k smallest *positive* eigenvalues λ_i for $i=1,2,\ldots,k$ are of interest. They lie in the middle of the spectrum of H, a region to which a straightforward application of the scaled $\mathcal{T}_m(\phi(t))$ often does not work well. This causes problems for applying Chebyshev filters. Fortunately, we can apply the squaring technique, noticing that the square of these eigenvalues are all real and lie in the left end of the spectrum of KM or MK. Moreover, KM and MK are diagonalizable by Lemma 2.1(d). In what follows, we restrict discussions on constructing Chebyshev filters for KM. Any development after minor modifications will work for MK as well.

Let $v \in \mathbb{C}^n$. It can be expressed in the basis $\{y_i\}_{i=1}^n$ as

$$v = \eta_1 y_1 + \eta_2 y_2 + \cdots + \eta_n y_n.$$

Suppose that we have three scalars $\alpha_0 < \alpha < \beta$ such that

$$\lambda_i^2 \in [\alpha_0, \beta] \text{ for } i = 1, 2, \dots, n, \text{ with } \lambda_k^2 < \alpha \text{ and } \lambda_{k+1}^2 \ge \alpha.$$
 (2.8)

In practice, initially α_0 , α , and β are likely only rough estimates and may violate (2.8), but they can be refined adaptively to satisfy (2.8) in the end. To amplify the components of v in y_i for i = 1, 2, ..., k and to suppress those in y_i for $i \ge k + 1$, we compute

$$\mathscr{T}_m(\phi(KM))v = \eta_1 \mathscr{T}_m(\phi(\lambda_1^2))y_1 + \eta_2 \mathscr{T}_m(\phi(\lambda_2^2))y_2 + \dots + \eta_n \mathscr{T}_m(\phi(\lambda_n^2))y_n.$$

Making $|\mathcal{T}_m(\phi(\lambda_i^2))| \le 1$ for i > k+1 while $|\mathcal{T}_m(\phi(\lambda_i^2))|$ for i = 1, 2, ..., k are (potentially much) larger than 1, $\mathcal{T}_m(\phi(KM))$ serves the purpose of filtering out the unwanted eigen-directions in v.

The same filtering idea works when v is replaced by a "block of vectors" $V \in \mathbb{C}^{n \times \ell}$ with a full column rank. Let

$$p_m(t) = \frac{\mathscr{T}_m(\phi(t))}{\mathscr{T}_m(\phi(\alpha_0))}, \quad \sigma_i = \frac{\mathscr{T}_{i-1}(\phi(\alpha_0))}{\mathscr{T}_i(\phi(\alpha_0))}.$$

It can be verified that

$$\sigma_1 = \omega/(\alpha_0 - \gamma),$$

$$\sigma_i = \left(\frac{2}{\sigma_1} - \sigma_{i-1}\right)^{-1} \quad \text{for } i = 2, 3, \dots,$$

where $\gamma = \frac{\alpha + \beta}{2}$ and $\omega = \frac{\beta - \alpha}{2}$. The scaled Chebyshev polynomials are defined by

$$p_0(t) = 1, \quad p_1(t) = (t - \gamma)\sigma_1/\omega,$$

 $p_i(t) = 2\sigma_i \frac{t - \gamma}{\omega} p_{i-1}(t) - \sigma_{i-1}\sigma_i p_{i-2}(t) \quad \text{for } i = 2, 3, \dots.$

```
\textbf{Algorithm 2.1} \ V_{\text{new}} = \texttt{Chebyshev\_filter}(V, m, \alpha, \beta, \alpha_0)
```

```
 \begin{array}{l} \mathbf{1} \ \ \omega = (\overline{\beta-\alpha})/2, \ \gamma = (\alpha+\beta)/2, \ \sigma_1 = \omega/(\alpha_0-\gamma), \ \tau = 2/\sigma_1. \\ \mathbf{2} \ \ V = (K(MV_{\mathrm{old}}) - \gamma V_{\mathrm{old}})\sigma_1/\omega. \\ \mathbf{3} \ \ \mathbf{for} \ \ i = 2:m \ \ \mathbf{do} \\ \qquad \qquad \sigma_i = 1/(\tau-\sigma_{i-1}), \ V_{\mathrm{new}} = 2(K(MV) - \gamma V)\sigma_i/\omega - \sigma_{i-1}\sigma_i V_{\mathrm{old}}. \\ \qquad V_{\mathrm{old}} = V, \ V = V_{\mathrm{new}}. \end{array}
```

Algorithm 2.1 displays the pseudo-code that implements the block version of the scaled Chebyshev filter for KM. It computes $V_{\text{new}} = p_m(KM)V$. The parameter α_0 in Algorithm 2.1 is mainly used for the purpose of scaling, and hence a crude estimate α_0 is usually sufficient. The lower bound α of unwanted eigenvalue interval can be estimated and refined during an iterative process of a subspace method with minimal or no extra computational cost. The upper bound β of eig(KM) needs to be supplied at the beginning. Though β can be bounded by $\|K\|\|M\|$ for any consistent matrix norm, it is often too large to make the filtering effective. In Section 3, we present a (much) tighter bound by constructing an estimator using the Lanczos biorthogonalization procedure.

2.2 The main algorithm

Let $V_{\text{new}} = p_m(KM)V$ be the output of the Chebyshev filter in Algorithm 2.1. Naturally the filtered columns V_{new} span an approximate invariant subspace of KM. Due to the equivalence of the eigenvalue problems (1.2) and (1.4a), the column space of V_{new} approximates the subspace spanned by the y-components of the eigenvectors of H associated with the designed eigenvalues λ_i , $i=1,2,\ldots,k$. To obtain the subspace spanned by the y-components, we can apply a Chebyshev filter on MK to get $U_{\text{new}} = p_m(MK)U$, the filtered columns in U_{new} span an approximate invariant subspace of MK, which also correspond to λ_i , $i=1,2,\ldots,k$. This works, but it introduces unnecessary cost. We can utilize the following lemma for a more cost effective method.

Lemma 2.2 Let \mathcal{Z} be an invariant subspace of H and let $Z = \begin{bmatrix} V \\ U \end{bmatrix}$ be the basis matrix of \mathcal{Z} with both V and U having n rows, then $\operatorname{span}(MV) = \operatorname{span}(U)$.



Lemma 2.2 follows from the structure of H. It suggests that if the columns of V span the y-component of an approximate invariant subspace of H, then the columns of U := MV should span the x-component of the same approximate invariant subspace. There are at least two advantages for computing U this way. First, it is economical because MV has to be computed in the later projection phase, and second, $W := V^H U = V^H MV$ is guaranteed to be symmetric positive definite (SPD). This SPD property is important in the later projection phase.

Now that we have $\{\text{span}(V), \text{span}(U)\}$ as a pair of approximate deflating subspaces of $\{K, M\}$, by [2], the best approximations to the first λ_i for i = 1, 2, ..., k within the pair of approximate deflating subspaces are the eigenvalues of ²

$$H_{\rm SR} = \begin{bmatrix} 0 & W_1^{-H} U^{H} K U W_1^{-1} \\ W_2^{-H} V^{H} M V W_2^{-1} & 0 \end{bmatrix}, \tag{2.9}$$

where W_1 and W_2 are two nonsingular factors of W as in $W = W_1^H W_2$. This reduced H_{SR} is closely related to the so-called *Rayleigh quotient pair* introduced in [29]. By [2, Theorem 2.9], we know that how W is factorized does not affect eventual approximations to the eigenvalues and the corresponding eigenvectors of H. Therefore we may take $W_1 = W_2 = R$, where $W = R^H R$ is its Cholesky decomposition. As a consequence.

$$H_{\rm SR} = \begin{bmatrix} 0 & G \\ I & 0 \end{bmatrix}$$
, where $G = W_1^{-H} U^{\rm H} K U W_1^{-1}$. (2.10)

Furthermore, if (μ^2, q) is an eigenpair of G, then $(\mu, \begin{bmatrix} \mu q \\ q \end{bmatrix})$ is an eigenpair of H_{SR} by Lemma 2.1(b). Therefore a corresponding approximate eigenpair of H can be taken as [2]

$$\left(\mu, \begin{bmatrix} \mu V R^{-1} q \\ U R^{-1} q \end{bmatrix}\right). \tag{2.11}$$

Algorithm 2.2 presents a piece of pseudo-code for a block Chebyshev-Davidson type method to compute the first n_{want} eigenpairs of LREP. In the algorithm, we apply the Chebyshev filter in Algorithm 2.1 to expand subspaces and seek the best approximate eigenpairs of H by computing the eigenpairs of the small size-2k matrix H_{SR} as in Eq. (2.10). For convenience, we identify this algorithm by BChebyDLR(ℓ), where ℓ is the block size. We prefer this block version because usually block type methods with relatively small block sizes are more competitive than non-block versions, especially when the desired eigenvalues have clusters or even multiples.

²This is only true if K is at least positive semi-definite. But later we show numerically that the eigenvalues of H_{SR} still provide very good approximations to λ_i for i = 1, 2, ..., k even when K is indefinite.



${\bf Algorithm~2.2}~{\tt BChebyDLR}(\ell)$

```
(i) V_0 = \text{randn}(n, \ell), \ \widetilde{V} = V_0, \ V = U = [], \ j = 1,
       k = 0 (k tracks the number of columns in V and U),
       s = 0 (s is the dimension of G),
       n_{\rm c} = 0 (n_{\rm c} is the number of converged eigenpairs).
(ii) Initial estimates: \beta, \alpha, \alpha_0, e.g., by Algorithm 3.2.
(iii) while (j \leq j_{\text{max}}) do
            \widetilde{V} = \mathtt{Chebyshev\_filter}(\widetilde{V}, m, \beta, \alpha, \alpha_0).
            Orthogonalize \widetilde{V} against U to obtain new \widetilde{V}.
   2
            Compute \widetilde{U} = M\widetilde{V}, \widetilde{V}^{H}\widetilde{U} = R^{H}R, and \widehat{U} = \widetilde{U}R^{-1}, \widehat{V} = \widetilde{V}R^{-1}.
   3
            Let \widehat{E} = K\widehat{U}, s = s + \text{size}(\widehat{U}, 2), k = s + n_c, V = [V, \widehat{V}] and U = [U, \widehat{U}].
   4
            \widehat{G} = U(:, n_{\rm c} + 1:k)^{\rm H} \widehat{E}, \ E = [E, \widehat{E}], \ {\rm and} \ G = \begin{bmatrix} G & \widehat{G} \\ \widehat{G}^{\rm H} & \widehat{U}^{\rm H} \widehat{E} \end{bmatrix}.
   5
            Compute the spectral decomposition: G = Q\Omega Q^{H} with the diagonal elements of
   6
            \Omega in the ascending order, and let k_{\text{old}} = k.
   7
            if (s > s_{\text{max}}) then restart: s = s_{\text{keep}}, k = n_{\text{c}} + s_{\text{keep}}.
            Update U_{(:,n_c+1:k)} = U_{(:,n_c+1:k_{\text{old}})}Q_{(:,1:s)}, V_{(:,n_c+1:k)} = V_{(:,n_c+1:k_{\text{old}})}Q_{(:,1:s)} and
   8
            E = EQ_{(:,1:s)}.
            Compute \ell Ritz pairs (\rho_i, \hat{z}_i) of G, with \hat{z}_i = \begin{bmatrix} \hat{y}_i \\ \hat{x}_i \end{bmatrix}, where \rho_i = \Omega^{\frac{1}{2}}_{(i,i)},
   9
            \hat{x}_i = U_{(:,n_c+i)} \text{ and } \hat{y}_i = \rho_i V_{(:,n_c+i)}.
            Test for convergence; t_c is the number of newly converged Ritz pairs; store the
 10
            converged Ritz pairs in the ascending order, and let n_c = n_c + t_c.
            if (n_c \ge n_{\text{want}}) then return.
 11
            if (t_c > 0), s_{old} = s then s = s - t_c, E_{(:,1:s)} = E_{(:,t_c+1:s_{old})} and
 12
            G_{(1:s,1:s)} = \Omega_{(t_c+1:s_{\text{old}},t_c+1:s_{\text{old}})}.
            \alpha = \text{median}(\text{diag}(\Omega)), \ \alpha_0 = \min(\text{diag}(\Omega)), \ \text{and update } \beta \text{ if necessary.}
 13
            V = V_{(:,n_c+1:n_c+\ell)}.
 14
            j = j + 1.
 15
```

A few remarks regarding Algorithm 2.2 are in order:

- 1. In our MATLAB implementation, initially we set V_0 to a random $n \times \ell$ matrix, e.g., as generated by MATLAB's randn, for simplicity. But if a better V_0 is known, we should use it.
- 2. Initially k = s, and then $k = s + n_c$ if some of the wanted Ritz pairs have converged, where n_c is the number of converged Ritz pairs (2.11). Converged Uq and Vq are stored in the first n_c columns of U and V, respectively.
- 3. According to Lemma 2.1(c), deflations are done by orthogonalizing the newly generated block \widetilde{V} against $U_{(:,1:n_c)}$ and \widetilde{U} against $V_{(:,1:n_c)}$. However, in Algorithm 2.2 the orthogonality between the columns of \widetilde{U} and those of $V_{(:,1:n_c)}$ is implied due to orthogonalization at step 2, according to Theorem 2.1 below.
- 4. Algorithm 3.2 in the next section provides an initial crude estimate of α , i.e., low bound of unwanted eigenvalues of KM, and the parameter α_0 for scaling. We will discuss this estimator in detail later. These estimates α_0 and α are updated at step 14, where median(diag(Ω)) is the MATLAB built-in function to return the



end

median value of the elements in the vector diag(Ω). Another choice for updating α was discussed in [30].

- 5. The estimated β by Algorithm 3.2 is a strict upper bound under a condition as listed in (3.2). But this condition is not easy to verify without expensive computations. When it is not satisfied, it is possible that the bound β returned from Algorithm 3.2 may be an underestimate, which can make the filter ineffective if the upper bound is not updated. In the numerical examples presented in Section 4 and many others we tried, β by Algorithm 3.2 is indeed a strict upper bound. However, to make the code robust, we need a way to address the case that β happens to be an underestimate. Fortunately this can be easily done by an adaptive strategy, as discussed at the end of Section 3. This adaptive strategy is applied in the numerical tests in Section 4.
- 6. At step 10, the convergence of a Ritz pair (ρ_i, \hat{z}_i) is tested by checking if its relative residual norm

$$\frac{\|H\hat{z}_i - \rho_i \hat{z}_i\|_1}{(\|H\|_1 + |\rho_i|)\|\hat{z}_i\|_1} \le \text{tol}, \tag{2.12}$$

where tol is a pre-set tolerance. At the same time, the swap procedure is to keep the converged Ritz values in the ascending order to make sure no wanted eigenvalues are missed. If K is indefinite, then pure imaginary Ritz values will appear. In this case, we sort them by the ascending order of their squares. Once we have new converged Ritz values, a purging procedure is used to remove the associated columns of E as well as the diagonal elements of G. This is simply done at step 12.

Computationally, the matrix spectral norm $\|\cdot\|_2$ is considerably more difficult to compute that the ℓ_1 matrix-operator norm $\|\cdot\|_1$. It is this consideration that leads us to use the ℓ_1 vector and matrix norm $\|\cdot\|_1$ in Eq. (2.12). Theoretically, the ℓ_2 vector and matrix-operator norm $\|\cdot\|_2$ are probably more preferable.

Theorem 2.1 Let U, V be two $n \times k$ matrices generated by Algorithm 2.2. Then $V^{H}U = V^{H}MV = I_{k}$.

Proof To distinguish the two U's and V's, we rewrite the assignments at step 4 as $U = [U_{\text{old}}, \widehat{U}]$ and $V = [V_{\text{old}}, \widehat{V}]$. During the first sweep of the **while** loop, $V^H U = I_\ell$ at step 4 because of how \widehat{U} and \widehat{V} are defined at step 3. Therefore $V^H_{\text{old}}U_{\text{old}} = I_{\widetilde{k}}$ is ensured from the previous sweep, where $\widetilde{k} = k - \ell$. Since $\widehat{V}^H \widehat{U} = I_\ell$ (because of step 3), and $\widehat{V}^H U_{\text{old}} = 0$ (because of step 2 and step 3), we have

$$V_{\mathrm{old}}^{\mathrm{H}}\widehat{U} = V_{\mathrm{old}}^{\mathrm{H}}M\widehat{V} = V_{\mathrm{old}}^{\mathrm{H}}M^{\mathrm{H}}\widehat{V} = U_{\mathrm{old}}^{\mathrm{H}}\widehat{V} = 0,$$

and therefore

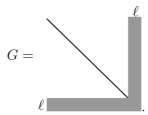
$$V^{\mathrm{H}}U = \begin{bmatrix} V_{\mathrm{old}}^{\mathrm{H}}U_{\mathrm{old}} & V_{\mathrm{old}}^{\mathrm{H}}\widehat{U} \\ \widehat{V}^{\mathrm{H}}U_{\mathrm{old}} & \widehat{V}^{\mathrm{H}}\widehat{U} \end{bmatrix} = I_{k},$$

as expected.



П

The eigenpairs of H_{SR} in Eq. (2.10) are obtained by computing the eigenpairs of G, where G at every step takes the following form:



This form is preserved in the algorithm when refining the basis matrices U and V at step 8. The new basis matrices UQ and VQ are reassigned to U and V. Refinement of basis matrices usually is applied in the thick restart method [12, 28] when the dimension of subspace exceeds the pre-set maximum dimension. Here, we refine the basis matrices U and V at every step. Although extra costs are incurred, eigenvectors often converge in fewer iterations [18, 22, 33]. Furthermore, the restart is easily executed by resetting the integers S and S when the size of S exceeds S max. The restart at step 7 is to keep the size of S and S under control. There are many ways to specify S and S we simply choose S max = S and S want and S we simply choose S max = S mush and S when S mush S mus

2.3 Convergence analysis for λ_1

A convergence analysis of the Chebyshev-Davidson method for the Hermitian eigenvalue problem was given in [30, 33]. Here, we analyze the convergence rate of Algorithm 2.2 per iteration step. Using the technique in [14] for estimating the accuracy of the smallest Ritz eigenvalue, we obtain an estimate in Theorem 2.2 below. In fact, Theorem 2.2 after minor modifications also holds for the Chebyshev-Davidson method for the Hermitian eigenvalue problem.

Suppose $\begin{bmatrix} v \\ u \end{bmatrix}$ is an approximate eigenvector corresponding to the first eigenvalue λ_1 of H, where u = Mv. We define

$$\rho(u, v) = \frac{u^{\mathrm{H}} K u}{u^{\mathrm{H}} v}.$$

Basing on the claim we made about the eigenvalues of Eq. (2.9) there, we can show that $\sqrt{\rho(u,v)}$ is the best approximation of λ_1 within the pair $\{u,v\}$. For proving Theorem 2.2, we need the following lemma that says $\rho(u,v)$ is an upper bound for the smallest Ritz value from $\{\text{span}(U), \text{span}(V)\}$.

Lemma 2.3 Let $U \in \mathbb{C}^{n \times k}$, $V \in \mathbb{C}^{n \times k}$ and ρ_i for $1 \leq i \leq k$ be computed in Algorithm 2.2, and let $u \in \operatorname{span}(U)$, $v \in \operatorname{span}(V)$, u = Mv and $\mu^2 = \min_{1 \leq i \leq k} \rho_i^2$. Then $\rho(u, v) \geq \mu^2$.



Proof As we know in Algorithm 2.2, $\rho_i^2 = \Omega_{(i,i)}$ for $1 \le i \le k$, where the diagonal elements of Ω are the eigenvalues of G. Let $v = V\tilde{v}$, where $\tilde{v} \in \mathbb{C}^k$. We have,

$$\begin{split} \mu^2 &= \min_{1 \leq i \leq k} \rho_i^2 = \min_{g \in \mathbb{C}^k} \frac{g^{\mathrm{H}} G g}{g^{\mathrm{H}} g} = \min_{g \in \mathbb{C}^k} \frac{g^{\mathrm{H}} V^{\mathrm{H}} M K M V g}{g^{\mathrm{H}} g}, \\ \rho(u, v) &= \frac{u^{\mathrm{H}} K u}{u^{\mathrm{H}} v} = \frac{v^{\mathrm{H}} M K M v}{v^{\mathrm{H}} M v} \\ &= \frac{\tilde{v}^{\mathrm{H}} V^{\mathrm{H}} M K M V \tilde{v}}{\tilde{v}^{\mathrm{H}} V^{\mathrm{H}} M V \tilde{v}} \\ &= \frac{\tilde{v}^{\mathrm{H}} V^{\mathrm{H}} M K M V \tilde{v}}{\tilde{v}^{\mathrm{H}} \tilde{v}^{\mathrm{H}} \tilde{v}}. \end{split}$$

The last equality holds because of $V^H M V = I_k$ by Theorem 2.1. Hence, $\rho(u, v) \ge \mu^2$.

Theorem 2.2 Let $(\mu_j, \begin{bmatrix} v_j \\ u_j \end{bmatrix})$ be from the j-th iteration of Algorithm 2.2, which is

the j-th approximation of the exact eigenvalue λ_1 and eigenvector $z_1 = \begin{bmatrix} \lambda_1 y_1 \\ x_1 \end{bmatrix}$, where y_1 and x_1 are defined by Eq. (2.6). Let α_{j+1} and β_{j+1} be the parameters α and β used in the (j+1)-st Chebyshev filter step, and α the degree of the Chebyshev polynomial. Define α (t), similarly to Eq. (2.3), as

$$\phi(t) = \frac{2t - (\alpha_{j+1} + \beta_{j+1})}{\beta_{j+1} - \alpha_{j+1}}.$$

Then

$$\frac{\mu_{j+1}^2 - \lambda_1^2}{\mu_j^2 - \lambda_1^2} \le \left(\frac{1-\xi}{1+\xi}\right)^2 + O(\delta_j^2),\tag{2.13}$$

where

$$\xi = \frac{\mathscr{T}_m(\phi(\lambda_1^2)) - \mathscr{T}_m(\phi(\lambda_{\diamond}^2))}{\mathscr{T}_m(\phi(\lambda_1^2)) - \mathscr{T}_m(\phi(\lambda_{*}^2))} \quad with$$

$$\mathscr{T}_m(\phi(\lambda_{*}^2)) = \min_{i \neq 1} \mathscr{T}_m(\phi(\lambda_i^2)), \quad \mathscr{T}_m(\phi(\lambda_{\diamond}^2)) = \max_{i \neq 1} \mathscr{T}_m(\phi(\lambda_i^2)),$$

and $\delta_j \to 0$, where $\delta_j = \sin \angle_M(v_j, y_1)$ is the sine of the angle between v_j to y_1 in the M-inner product.

Proof Using the same notations as before, we know $KM = Y\Lambda^2 X^H$ with $Y = [y_1, y_2, ..., y_n], X = [x_1, x_2, ..., x_n]$, and

$$v_j = \mu_j V q_1, \quad u_j = U q_1,$$
 (2.14)

where q_1 is the eigenvector of G corresponding to μ_j and $||q_1||_2 = 1$.



Let $\|\cdot\|_M$ denote the norm induced by the *M*-inner product. Since $My_1 = x_1$ and $y_1^H x_1 = 1$, we have $\|y_1\|_M = 1$. Set

$$\tilde{v}_j = \frac{v_j}{\|v_j\|_M} = \frac{\mu_j V q_1}{\mu_j} = V q_1.$$

Then \tilde{v}_j has the decomposition $\tilde{v}_j = \gamma_j y_1 + \delta_j s_j$, where $s_j \in \text{span}\{y_2, \dots, y_n\}$, $\|s_j\|_M = 1$ and $\gamma_j^2 + \delta_j^2 = 1$. As stated in [30, 33], $\delta_j \to 0$ at least as fast as by the power method.

By Eq. (2.14), we have $u_j = M\tilde{v}_j = \gamma_j M y_1 + \delta_j M s_j$, and it is not hard to see that $y_1^H M K M s_j = 0$ and $y_1^H M s_j = 0$. Therefore,

$$\mu_{j}^{2} = q_{1}^{H}Gq_{1} = u_{j}^{H}Ku_{j}$$

$$= (\gamma_{j}My_{1} + \delta_{j}Ms_{j})^{H}K(\gamma_{j}My_{1} + \delta_{j}Ms_{j})$$

$$= (\gamma_{j}My_{1} + \delta_{j}Ms_{j})^{H}(\lambda_{1}^{2}\gamma_{j}y_{1} + \delta_{j}KMs_{j})$$

$$= \gamma_{j}^{2}\lambda_{1}^{2} + \delta_{j}^{2}s_{j}^{H}MKMs_{j}$$

$$= \lambda_{1}^{2} + \delta_{i}^{2}(s_{j}^{H}MKMs_{j} - \lambda_{1}^{2}). \tag{2.15}$$

Let \hat{v}_{j+1} be a linear combination of \tilde{v}_j and $\mathcal{T}_m(\phi(KM))\tilde{v}_j$, i.e., $\hat{v}_{j+1} = \tilde{v}_j - \tau \mathcal{T}_m(\phi(KM))\tilde{v}_j$ for some scalar τ , and let $\hat{u}_{j+1} = M\hat{v}_{j+1}$. Then

$$\hat{v}_{j+1} = \gamma_j [1 - \tau \mathcal{T}_m(\phi(\lambda_1^2)))] y_1 + \delta_j [I - \tau \mathcal{T}_m(\phi(KM))] s_j,$$

$$\hat{u}_{j+1} = \gamma_j [1 - \tau \mathcal{T}_m(\phi(\lambda_1^2))] M y_1 + \delta_j M [I - \tau \mathcal{T}_m(\phi(KM))] s_j.$$

The best approximation of λ_1 within the pair $\{\hat{u}_{j+1}, \hat{v}_{j+1}\}$ is $\sqrt{\rho(\hat{u}_{j+1}, \hat{v}_{j+1})}$. Substituting $t_1 = 1 - \tau \mathcal{T}_m(\phi(\lambda_1^2))$ and $\widetilde{T} = I - \tau \mathcal{T}_m(\phi(KM))$ in $\rho(\hat{u}_{j+1}, \hat{v}_{j+1})$, we get

$$\begin{split} \rho(\hat{u}_{j+1}, \hat{v}_{j+1}) &= \frac{\hat{u}_{j+1}^{H} K \hat{u}_{j+1}}{\hat{u}_{j+1}^{H} \hat{v}_{j+1}} \\ &= \frac{\gamma_{j}^{2} t_{1}^{2} \lambda_{1}^{2} + \delta_{j}^{2} s_{j}^{H} \widetilde{T}^{H} M K M \widetilde{T} s_{j}}{\gamma_{j}^{2} t_{1}^{2} + \delta_{j}^{2} s_{j}^{H} \widetilde{T}^{H} M \widetilde{T} s_{j}} \\ &= \lambda_{1}^{2} + \frac{\delta_{j}^{2} s_{j}^{H} \widetilde{T}^{H} (M K M - \lambda_{1}^{2} M) \widetilde{T} s_{j}}{\gamma_{j}^{2} t_{1}^{2} + \delta_{j}^{2} s_{j}^{H} \widetilde{T}^{H} M \widetilde{T} s_{j}} \\ &= \lambda_{1}^{2} + \frac{(\delta_{j}^{2} / t_{1}^{2}) s_{j}^{H} \widetilde{T}^{H} (M K M - \lambda_{1}^{2} M) \widetilde{T} s_{j}}{1 + (\delta_{j}^{2} / t_{1}^{2}) (s_{j}^{H} \widetilde{T}^{H} M \widetilde{T} s_{j} - t_{1}^{2})} \\ &= \lambda_{1}^{2} + (\delta_{j}^{2} / t_{1}^{2}) s_{j}^{H} \widetilde{T}^{H} (M K M - \lambda_{1}^{2} M) \widetilde{T} s_{j} + O(\delta_{j}^{4}). \end{split}$$

Let μ_{j+1} be the (j+1)-st approximation of λ_1 computed by Algorithm 2.2. Then by Lemma 2.3, we have $\mu_{j+1}^2 \leq \rho(\hat{u}_{j+1}, \hat{v}_{j+1})$. Set

$$\widehat{T} = I - \tau \, \mathscr{T}_m(\phi(\Lambda^2)), \quad t_\tau^2 = \max_{i \neq 1} \left[1 - \tau \, \mathscr{T}_m(\phi(\lambda_i^2)) \right]^2.$$



We have

$$\begin{split} \mu_{j+1}^2 - \lambda_1^2 &\leq \rho(\hat{u}_{j+1}, \hat{v}_{j+1}) - \lambda_1^2 \\ &= (\delta_j^2/t_1^2) s_j^H \widetilde{T}^H (MKM - \lambda_1^2 M) \widetilde{T} s_j + O(\delta_j^4) \\ &= (\delta_j^2/t_1^2) s_j^H X \widehat{T} (\Lambda^2 - \lambda_1^2 I) \widehat{T} X^H s_j + O(\delta_j^4) \\ &= (\delta_j^2/t_1^2) s_j^H X (\Lambda^2 - \lambda_1^2 I)^{\frac{1}{2}} \widehat{T} \widehat{T} (\Lambda^2 - \lambda_1^2 I)^{\frac{1}{2}} X^H s_j + O(\delta_j^4) \\ &\leq (t_\tau^2/t_1^2) \delta_j^2 s_j^H X (\Lambda^2 - \lambda_1^2 I)^{\frac{1}{2}} (\Lambda^2 - \lambda_1^2 I)^{\frac{1}{2}} X^H s_j + O(\delta_j^4) \\ &= (t_\tau^2/t_1^2) \delta_j^2 s_j^H (MKM - \lambda_1^2 M) s_j + O(\delta_j^4) \\ &= (t_\tau^2/t_1^2) (\mu_i^2 - \lambda_1^2) + O(\delta_j^4). \end{split}$$
 (by Eq. 2.15)

Therefore, by Eq. (2.15), we have $\mu_i^2 - \lambda_1^2 = O(\delta_i^2)$ and

$$\frac{\mu_{j+1}^2 - \lambda_1^2}{\mu_j^2 - \lambda_1^2} \le \min_{\tau} \frac{t_{\tau}^2}{t_1^2} + O(\delta_j^2).$$

From the definition of t_{τ} , we know that

$$t_{\tau}^2 = \max \left\{ \left[1 - \tau \mathcal{T}_m(\phi(\lambda_{\diamond}^2)) \right]^2, \left[1 - \tau \mathcal{T}_m(\phi(\lambda_{*}^2)) \right]^2 \right\},$$

where
$$\mathscr{T}_m(\phi(\lambda_*^2)) = \min_{i \neq 1} \mathscr{T}_m(\phi(\lambda_i^2))$$
 and $\mathscr{T}_m(\phi(\lambda_\diamond^2)) = \max_{i \neq 1} \mathscr{T}_m(\phi(\lambda_i^2))$.

We first consider the case where $\mathscr{T}_m(\phi(\lambda_{\diamond}^2)) + \mathscr{T}_m(\phi(\lambda_{*}^2)) \geq 0$. Under this additional condition,

$$t_{\tau}^2 = \begin{cases} \left[1 - \tau \mathcal{T}_m(\phi(\lambda_{\diamond}^2))\right]^2, \text{ for } \tau \geq 2 / [\mathcal{T}_m(\phi(\lambda_{\diamond}^2)) + \mathcal{T}_m(\phi(\lambda_{*}^2))] \text{ or } \tau \leq 0, \\ \left[1 - \tau \mathcal{T}_m(\phi(\lambda_{*}^2))\right]^2, \text{ for } 0 \leq \tau \leq 2 / [\mathcal{T}_m(\phi(\lambda_{\diamond}^2)) + \mathcal{T}_m(\phi(\lambda_{*}^2))]. \end{cases}$$

For
$$0 \le \tau \le 2/[\mathscr{T}_m(\phi(\lambda_\diamond^2)) + \mathscr{T}_m(\phi(\lambda_*^2))]$$
,

$$\frac{\mathrm{d}(t_\tau/t_1)^2}{\mathrm{d}\tau} = \frac{2\left[\mathcal{T}_m(\phi(\lambda_1^2)) - \mathcal{T}_m(\phi(\lambda_*^2))\right]\left[\tau\mathcal{T}_m(\phi(\lambda_1^2)) - 1\right]\left[\tau\mathcal{T}_m(\phi(\lambda_*^2)) - 1\right]}{\left[1 - \tau\mathcal{T}_m(\phi(\lambda_1^2))\right]^4}.$$

Therefore, if $\mathcal{T}_m(\phi(\lambda_1^2)) > 0$, then

$$\begin{cases} \frac{\mathrm{d}(t_{\tau}/t_1)^2}{\mathrm{d}\tau} \geq 0, \text{ for } 0 \leq \tau \leq 1/[\mathcal{T}_m(\phi(\lambda_1^2))], \\ \frac{\mathrm{d}(t_{\tau}/t_1)^2}{\mathrm{d}\tau} \leq 0, \text{ for } 1/[\mathcal{T}_m(\phi(\lambda_1^2))] \leq \tau \leq 2/[\mathcal{T}_m(\phi(\lambda_\diamond^2)) + \mathcal{T}_m(\phi(\lambda_*^2))]. \end{cases}$$

Meanwhile, for $\tau \ge 2/[\mathscr{T}_m(\phi(\lambda_*^2)) + \mathscr{T}_m(\phi(\lambda_*^2))]$ or $\tau \le 0$, it is not hard to prove

$$\begin{cases} \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \geq 0, \text{ for } \tau \leq 0, \\ \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \geq 0, \text{ for } \tau \geq 2/[\mathscr{T}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathscr{T}_{m}(\phi(\lambda_{*}^{2}))]. \end{cases}$$



When $\mathcal{T}_m(\phi(\lambda_1^2)) < 0$, we have

$$\begin{cases} \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \leq 0, \text{ for } 0 \leq \tau \leq 2/[\mathcal{T}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathcal{T}_{m}(\phi(\lambda_{*}^{2}))], \\ \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \leq 0, \text{ for } 1/\mathcal{T}_{m}(\phi(\lambda_{1}^{2}))] \leq \tau \leq 0, \\ \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \geq 0, \text{ for } \tau \leq 1/\mathcal{T}_{m}(\phi(\lambda_{1}^{2}))], \\ \frac{\mathrm{d}(t_{\tau}/t_{1})^{2}}{\mathrm{d}\tau} \geq 0, \text{ for } \tau \geq 2/[\mathcal{T}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathcal{T}_{m}(\phi(\lambda_{*}^{2}))]. \end{cases}$$

Therefore

$$\min_{\tau} \frac{t_{\tau}^{2}}{t_{1}^{2}} = \frac{\left(1 - \frac{2}{\mathcal{I}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathcal{I}_{m}(\phi(\lambda_{*}^{2}))} \mathcal{I}_{m}(\phi(\lambda_{\diamond}^{2}))\right)^{2}}{\left(1 - \frac{2}{\mathcal{I}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathcal{I}_{m}(\phi(\lambda_{*}^{2}))} \mathcal{I}_{m}(\phi(\lambda_{1}^{2}))\right)^{2}} \\
= \frac{\left(\mathcal{I}_{m}(\phi(\lambda_{\diamond}^{2})) - \mathcal{I}_{m}(\phi(\lambda_{*}^{2}))\right)^{2}}{\left(\mathcal{I}_{m}(\phi(\lambda_{\diamond}^{2})) + \mathcal{I}_{m}(\phi(\lambda_{*}^{2})) - 2\mathcal{I}_{m}(\phi(\lambda_{1}^{2}))\right)^{2}} \\
= \left(\frac{1 - \xi}{1 + \xi}\right)^{2},$$

where $\xi = [\mathscr{T}_m(\phi(\lambda_1^2)) - \mathscr{T}_m(\phi(\lambda_\diamond^2))]/[\mathscr{T}_m(\phi(\lambda_1^2)) - \mathscr{T}_m(\phi(\lambda_*^2))].$ The remaining case where $\mathscr{T}_m(\phi(\lambda_\diamond^2)) + \mathscr{T}_m(\phi(\lambda_*^2)) \leq 0$ can be addressed in a

The remaining case where $\mathscr{T}_m(\phi(\lambda_\diamond^2)) + \mathscr{T}_m(\phi(\lambda_*^2)) \leq 0$ can be addressed in a similar way to derive the same result.

Once the convergence rate for λ_1 is established, according to the remark 3 for Algorithm 2.2, the convergence analysis for other eigenvalues $\lambda_2, \dots, \lambda_k$ can be addressed using the argument of deflation. For example, with a similar argument, the convergence rate formula for λ_2 can be gotten as

$$\frac{\mu_{j+1}^2 - \lambda_2^2}{\mu_j^2 - \lambda_2^2} \le \left(\frac{1 - \xi}{1 + \xi}\right)^2 + O(\delta_j^2),\tag{2.16}$$

where

$$\xi = \frac{\mathscr{T}_m(\phi(\lambda_2^2)) - \mathscr{T}_m(\phi(\lambda_2^2))}{\mathscr{T}_m(\phi(\lambda_2^2)) - \mathscr{T}_m(\phi(\lambda_*^2))} \quad \text{with}$$

$$\mathscr{T}_m(\phi(\lambda_*^2)) = \min_{i \neq 1, 2} \mathscr{T}_m(\phi(\lambda_i^2)), \quad \mathscr{T}_m(\phi(\lambda_\diamond^2)) = \max_{i \neq 1, 2} \mathscr{T}_m(\phi(\lambda_i^2)),$$

and $\delta_j = \sin \angle_M(v_j, y_2)$.

Similar convergence estimate appears in the constant-shift stationary Richardson method for computing the largest eigenvalue of generalized eigenvalue problems [10]. More convergence estimates for the generalized Davidson method can be found in [15, 16].



3 Upper bound for eig(KM)

Theoretically, $\|K\|_1\|M\|_1$ and $\|K\|_\infty\|M\|_\infty$ are upper bounds of λ^2 and they are easily computable, where λ is an eigenvalue of H, but they are usually too large for the Chebyshev filter to be effective. A sharper upper bound is needed. The k-step Lanczos method for computing a sharp upper bound of the spectrum of a symmetric matrix is proposed in [32]. In this section, we propose a similar technique to compute an upper bound of the spectrum of KM.

Lemma 3.1 ([5, Theorem 1]) Let Ω be an $n \times n$ real diagonal matrix, D a $k \times k$ diagonal matrix with diagonal elements $d_1 \leq d_2 \leq \cdots \leq d_k$, and let Q be an $n \times k$ matrix having full column rank, where $1 \leq k \leq n$. Then there exist k diagonal elements $\mu_{i_1} \leq \mu_{i_2} \leq \cdots \leq \mu_{i_k}$ of Ω such that

$$\|\Omega Q - QD\|_2 \ge \max_{1 \le i \le k} |d_j - \mu_{i_j}| \sigma_{\min}(Q),$$

where $\sigma_{\min}(Q)$ is the smallest singular value of Q.

To state the next theorem, we introduce, in addition to the notations in Lemma 2.1, $\lambda_i(KM)$ for $1 \le i \le n$ for the eigenvalues of KM in the ascending order, and

$$\lambda_{\max}(KM) := \max_{i} \lambda_{i}(KM) = \lambda_{n}(KM) = \lambda_{n}^{2}.$$

Theorem 3.1 Let U and V be two $n \times k$ matrices with full column rank, $H_K = U^H K U$, and $H_M = V^H M V$, where $1 \le k \le n$, and let $\theta_1 \le \theta_2 \le \cdots \le \theta_k$ be the eigenvalues of $H_K H_M$. Then there exist k eigenvalues $\lambda_{i_1}^2 \le \lambda_{i_2}^2 \le \cdots \le \lambda_{i_k}^2$ of KM such that

$$\max_{1 \le j \le k} |\theta_j - \lambda_{i_j}^2| \le \sqrt{\|M\|_2 \|H_M^{-1}\|_2} \|KMV - VH_K H_M\|_2$$
 (3.1a)

$$\leq \sqrt{\|M\|_1\|H_M^{-1}\|_1} \|KMV - VH_KH_M\|_2.$$
 (3.1b)

If, in addition,

$$|\lambda_{\max}(KM) - \lambda_{\max}(H_K H_M)| = \min_{1 \le i \le n} |\lambda_i(KM) - \lambda_{\max}(H_K H_M)|, \qquad (3.2)$$

where $\lambda_{\max}(H_K H_M) = \theta_k =: \theta_{\max}$, then

$$\lambda_n^2 \le \theta_{\text{max}} + \sqrt{\|M\|_2 \|H_M^{-1}\|_2} \|KMV - VH_KH_M\|_2$$
 (3.3a)

$$\leq \theta_{\text{max}} + \sqrt{\|M\|_1 \|H_M^{-1}\|_1} \|KMV - VH_K H_M\|_2. \tag{3.3b}$$

Proof The inequalities in Eq. (3.3) are consequences of Eq. (3.1) by Eq. (3.2). So we will prove (3.1) only. By Lemma 2.1(a), we have $K = Y\Lambda^2Y^H$,



 $M = XX^{\mathrm{H}}, H_K = \widetilde{Y}\Theta\widetilde{Y}^{\mathrm{H}}, \text{ and } H_M = \widetilde{X}\widetilde{X}^{\mathrm{H}}, \text{ where } \Theta = \mathrm{diag}(\theta_1, \theta_2, \dots, \theta_k).$ We have

$$\begin{split} \|KMV - VH_KH_M\|_2 &= \|Y\Lambda^2X^{H}V - VH_KH_M\|_2 \\ &= \|Y(\Lambda^2X^{H}V - X^{H}VH_KH_M)\|_2 \\ &= \|Y(\Lambda^2X^{H}V\widetilde{Y} - X^{H}V\widetilde{Y}\Theta)\widetilde{X}^{H}\|_2 \\ &\geq \frac{\|\Lambda^2X^{H}V\widetilde{Y} - X^{H}V\widetilde{Y}\Theta\|_2}{\|X^{H}\|_2\|\widetilde{Y}\|_2}. \end{split}$$

According to Lemma 3.1, there exist $\lambda_{i_1}^2 \leq \lambda_{i_2}^2 \leq \cdots \leq \lambda_{i_k}^2$ such that

$$\|\Lambda^2 X^{\mathrm{H}} V \widetilde{Y} - X^{\mathrm{H}} V \widetilde{Y} \Theta\|_2 \geq \sigma_{\min} \big(X^{\mathrm{H}} V \widetilde{Y} \big) \max_{1 \leq j \leq k} |\theta_j - \lambda_{i_j}^2| = \max_{1 \leq j \leq k} |\theta_j - \lambda_{i_j}^2|.$$

The last equality holds because of $\sigma_{\min}(X^HV\widetilde{Y}) = \sqrt{\lambda_{\min}(\widetilde{Y}^HV^HXX^HV\widetilde{Y})} = 1$. Therefore

$$||KMV - VH_KH_M||_2 \ge \frac{1}{||X^H||_2||\widetilde{Y}||_2} \max_{1 \le j \le k} |\theta_j - \lambda_{i_j}^2|.$$

Now use

$$\|X^{\mathbf{H}}\|_{2} = \sqrt{\|M\|_{2}} \le \sqrt{\|M\|_{1}}, \quad \|\widetilde{Y}\|_{2} = \sqrt{\|H_{M}^{-1}\|_{2}} \le \sqrt{\|H_{M}^{-1}\|_{1}}$$
 to conclude the proof of Eq. (3.1).

The inequality (3.3) provides upper bounds on λ_n^2 . If the eigenvector associated with θ_{max} is also available, we have the following theorem for an improved bound.

Theorem 3.2 Under the conditions of Theorem 3.1, including (3.2). Let w be an eigenvector corresponding to the largest eigenvalue θ_{max} of $H_K H_M$, normalized such that $w^H H_M w = 1$. Then

$$\lambda_n^2 \le \theta_{\text{max}} + \sqrt{\|M\|_2} \|KMVw - VH_KH_Mw\|_2$$
 (3.4a)

$$\leq \theta_{\max} + \sqrt{\|M\|_1} \|KMVw - VH_KH_Mw\|_2. \tag{3.4b}$$

Proof We have

$$||KMVw - VH_KH_Mw||_2 = ||KMVw - \theta_{\max}Vw||_2$$

$$= ||Y\Lambda^2X^HVw - \theta_{\max}Vw||_2$$

$$= ||Y(\Lambda^2 - \theta_{\max}I_n)X^HVw||_2$$

$$\geq \frac{||(\Lambda^2 - \theta_{\max}I_n)X^HVw||_2}{||X^H||_2}$$

$$\geq \frac{|w^HV^HXX^HVw|^{1/2}}{||X^H||_2}|\lambda_n^2 - \theta_{\max}| \quad \text{(by Eq. 3.2)}$$

$$\geq \frac{|w^HV^HMVw|^{1/2}}{||X^H||_2}|\lambda_n^2 - \theta_{\max}|.$$



Now use $||X^{H}||_2 = \sqrt{||M||_2} \le \sqrt{||M||_1}$ to conclude the proof.

Algorithm 3.1 Lanczos biorthogonalization procedure

$$\begin{array}{l} \mathbf{1} \ \ v = \mathrm{randn}(n,1), \ u = Mv, \ b = \sqrt{u^{\mathrm{H}}v}, \ u_1 = u/b, \ v_1 = v/b, \ b_0 = 0. \\ \mathbf{2} \ \ \mathbf{for} \ \ i = 1:k \ \ \mathbf{do} \\ a_i = u_i^{\mathrm{H}} K u_i. \\ \tilde{v} = K u_i - a_i v_i - b_{i-1} v_{i-1}, \ \tilde{u} = M \tilde{v}. \\ b_i = \sqrt{\tilde{u}^{\mathrm{H}} \tilde{v}}. \\ u_{i+1} = \tilde{u}/b_i, \ v_{i+1} = \tilde{v}/b_i. \\ \mathbf{end} \end{array}$$

The bounds (3.4) can be significantly simplified if the following Lanczos biorthogonalization procedure [23, subsection 3.1] is applied to compute θ_{max} . Suppose that Algorithm 3.1 successfully runs to its completion, i.e., $b_i \neq 0$ for all i. Denote $U_k = [u_1, u_2, \dots, u_k], V_k = [v_1, v_2, \dots, v_k]$, and

$$T_{k} = \begin{bmatrix} a_{1} & b_{1} \\ b_{1} & a_{2} & b_{2} \\ & \ddots & \ddots & \ddots \\ & b_{k-2} & a_{k-1} & b_{k-1} \\ & & b_{k-1} & a_{k} \end{bmatrix}.$$
(3.5)

Then

$$KU_k = V_k T_k + b_k v_{k+1} e_k^{\mathrm{T}}, \quad MV_k = U_k,$$

and also $U_k^H V_k = I_k$, $U_k^H K U_k = T_k$, and $V_k^H M V_k = I_k$.

Corollary 3.1 In Algorithm 3.1, let $T_k w = \theta_{\text{max}} w$, $||w||_2 = 1$, where θ_{max} is the largest eigenvalue of T_k . If Eq. (3.2) holds, then

$$\lambda_n^2 \le \theta_{\max} + \sqrt{\|M\|_2} \|b_k(e_k^T w)v_{k+1}\|_2$$
 (3.6a)

$$\leq \theta_{\max} + \sqrt{\|M\|_1} \|b_k(e_k^{\mathsf{T}}w)v_{k+1}\|_2.$$
 (3.6b)

Algorithm 3.2 An estimator for the bounds used in the first filtering step

Input: K, M and the number of the Lanczos step k_p .

Output: β , α and α_0 as discussed in subsection 2.1.

- 1 Generate V_k , $T_k v_{k+1}$ and b_k by Algorithm 3.1 with $k = k_p$.
- 2 Compute eigen-decomposition $T_k = Z^H \Theta Z$, and let (θ_{max}, w) be the largest eigenpair of T_k , where $||w||_2 = 1$.
- 3 Compute $\beta = \theta_{\max} + \sqrt{\|M\|_1} \|b_k(e_k^T w)v_{k+1}\|_2$ by (3.6b), and let $\alpha = \text{median}(\Theta)$, $\alpha_0 = \min(\Theta)$.

With these preparations, we propose Algorithm 3.2 to compute an upper bound for eig(KM). To test the effectiveness of Algorithm 3.2, we compose 6 test problems, listed as TEST 1 to TEST 6 in Table 1. The K- and M-matrices of TEST 1 to TEST 3 come from the linear response analysis for Na2, Na4, and silane (SiH4) compound, respectively. These matrices are generated by the turboTDDFT code in QUANTUM ESPRESSO — an electronic structure calculation code that implements density



Table 1	Test problems	

Problem	TEST 1	TEST 2	TEST 3	TEST 4	TEST 5	TEST 6
n	1862	2834	5660	5832	5743	74752
K	Na2	Na4	\mathtt{SiH}_4	Na5	SiNa	\mathtt{SiO}_2
M	Na2	Na4	\mathtt{SiH}_4	fv1	fv2	finan512

functional theory (DFT) using plane-waves as the basis set and pseudopotentials [9]. For Na2, Na4 and SiH4, the matrices K and M are symmetric positive definite. TEST 4 to TEST 6, artificially constructed, consist of matrices from the University of Florida Sparse Matrix Collection [7] to give K and M. In the case when the two matrices from the collection have different dimensions, we extract out the leading principal submatrix of the larger one to give K and M of equal size. For TEST 4 and TEST 6, the M-matrices are SPD, but the K-matrices are indefinite.

We apply the built-in function eigs of MATLAB with default tolerance tol = 2.22×10^{-16} to compute the largest eigenvalue λ_n of associated H to the "exact" eigenvalue bound λ_n^2 . These test problems are also used in the next section to test BChebyDLR(ℓ) (Algorithm 2.2).

Figure 1 plots the upper bounds by Algorithm 3.2 as k_p varies for the six tests, where the red dotted lines are for the "exact" λ_n^2 , and the blue solid lines are the upper bounds by Algorithm 3.2. It demonstrates that Algorithm 3.2 is rather efficient and produces a fairly tight upper bound on λ_n^2 in a few Lanczos steps, such as within 10 steps.

The inequality (3.6b) is proved under the condition of Eq. (3.2) which, nevertheless, may fail for the first few Lanczos steps. When it does, it may underestimate λ_n^2

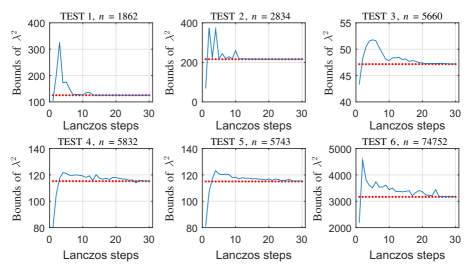


Fig. 1 Behavior of the upper bound estimator. The red dotted lines are for the "exact" λ_n^2 , and the blue solid lines are the upper bounds by Algorithm 3.2



as shown in Fig. 1. In practice such an underestimation does not cause any problem and can be overcome by using an adaptive strategy within Algorithm 2.2 to update the upper bound. Let ρ_{max} be the largest eigenvalue of G in Algorithm 2.2. For any nonzero $g \in \mathbb{C}^k$, where k is the dimension of G, we have

$$\rho_{\text{max}} = \max_{g \in \mathbb{C}^k} \frac{g^{\text{H}} G g}{g^{\text{H}} g} = \max_{g \in \mathbb{C}^k} \frac{g^{\text{H}} V^{\text{H}} M K M V g}{g^{\text{H}} g}$$

$$= \max_{g \in \mathbb{C}^k} \frac{g^{\text{H}} V^{\text{H}} X \Lambda^2 X^{\text{H}} V g}{g^{\text{H}} g} \qquad \text{(by Eq. 2.5)}$$

$$\leq \lambda_n^2 \max_{g \in \mathbb{C}^k} \frac{g^{\text{H}} V^{\text{H}} X X^{\text{H}} V g}{g^{\text{H}} g}$$

$$= \lambda_n^2. \qquad (3.7)$$

Let β be computed by Algorithm 3.2. Then by the sign of $\beta - \rho_{max}$, we find whether we need to update the upper bound. Namely, if $\rho_{max} > \beta$, then we update β as $\beta = \rho_{max}$. We demonstrate by numerical examples in the next section that this adaptive strategy upon integrating into BChebyDLR works very well.

4 Numerical results

We present some numerical results of Algorithm 2.2, denoted by BChebyDLR(ℓ), to compute the first n_{want} eigenpairs. Recall that the first few λ_i may be purely imaginary if K is indefinite. A computed eigenpair (ρ , \hat{z}) is considered as converged when its relative residual norm is below a user specified tol, which by default is set to 10^{-8} , i.e.,

$$\frac{\|H\hat{z} - \rho\hat{z}\|_1}{(\|H\|_1 + |\rho|)\|\hat{z}\|_1} \le \text{tol} = 10^{-8}.$$

The test matrices are TEST 1 to TEST 6 in Table 1. TEST 1 to TEST 3 are real LREPs from physics, and for the technical reason, the K- and M-matrices are dense and symmetric positive definite, whereas the matrices of TEST 4 to TEST 6 are sparse and symmetric with K-matrices being indefinite and M-matrices still being positive definite. We use these matrices to show the effectiveness of our proposed algorithm for eigenproblems of form (1.2), and this form includes LREP as a special case.

In all tests, we start with $V_0 = \text{randn}(n, \ell)$, where ℓ is the block size. The parameters n_{want} and the Chebyshev filter degree m are to be specified. The upper bound β is obtained by Algorithm 3.2 with $k_p = 10$, and $s_{\text{max}} = 2n_{\text{want}}$ when $n_{\text{want}} < 100$. But we use $s_{\text{max}} \le \text{ceil}(1.5n_{\text{want}})$ when $n_{\text{want}} \ge 100$. For restart, we use $s_{\text{keep}} = n_{\text{want}}$.

The hardware used for the numerical computations is a laptop with 8G memory, the CPU type is Intel core i5-3210M (a) 2.50GHz. We use MATLAB version 8.5 (R2015a).

Example 4.1 We first test BChebyDLR (ℓ) on the three problems from the linear response analysis for Na2, Na4 and SiH4, i.e., TEST 1 to TEST 3. Since both K and M



are positive definitive for these problems, we can compare BChebyDLR (ℓ) with the locally optimal block *preconditioned* 4-D CG method (LOBP4DCG) and the locally optimal block 4-D CG method (LOB4DCG) in [3].

For the LOBP4DCG method, we use the generic preconditioner

$$\Phi = \begin{bmatrix} 0 & M^{-1} \\ K^{-1} & 0 \end{bmatrix}. \tag{4.1}$$

The preconditioned search vectors q_i and p_i in [3] are computed by using the linear CG method [8]. Often very crude approximations of q_i and p_i are good enough. In this example, the linear CG iterations are set maximal 5 iterations. It is denoted by LOBP4DCG(5). The initial block size in LOBP4DCG(5) and LOB4DCG are chosen to be the same as $n_{\rm want}$. A deflation procedure is built to purge out converged eigenvectors. So the block size decreases as the algorithms progress. As for BChebyDLR, converged eigenvectors are also deflated from the iteration subspace, and new basis vectors need to be orthogonalized against the deflated eigenvectors.

We report the total number of matrix-vector products (denoted by "#mvp"), iteration number (denoted by "#iter"), and CPU time in seconds for BChebyDLR, LOB4DCG and LOBP4DCG(5) in Table 2.

We make a few comments on the two methods applied to these definite LREPs. First, for the three tests as well as other tests not reported here, BChebyDLR(ℓ) appears to be close to an order of magnitude faster than LOBP4DCG. However, we also observe that for some other problems not reported here, the LOBP4DCG can be an order of magnitude faster than BChebyDLR(ℓ). The main reason for this quite drastic difference is the conditioning of the KM-matrix. This may be similar to SVD calculations, e.g., squaring of a matrix makes the smallest singular values harder to converge to high accuracy. Our algorithm appears to be more sensitive to the conditioning of M. If M is well-conditioned, then generally BChebyDLR(ℓ) is much faster than LOBP4DCG; while for M that has a relatively large condition number, the LOBP4DCG is usually much more efficient than BChebyDLR(ℓ). Therefore, the two methods appear to be complementary to each other. Second, the effectiveness

Table 2 The number of matrix-vector products (#mvp), number of iterations (#iter), and CPU time in seconds for computing $n_{\text{want}} = 180$ eigenpairs. For BChebyDLR the filter degree used is m = 25, and the block size is $\ell = 15$. For LOB4DCG and LOBP4DCG (5) the initial block size is n_{want}

	BChebyDLR(ℓ)			LOB4DCG			LOBP4DCG(5)		
	#mvp	#iter	CPU	#mvp	#iter	CPU	#mvp	#iter	CPU
TEST 1	26520	34	49.9	40040	120	201.9	48600	61	235.9
TEST 2	71760	92	223.5	63712	812	462.8	56394	239	305.4
TEST 3	37440	48	295.5	109568	233	1028.5	161658	142	2960.2

The maximum subspace dimension used for BChebyDLR is 233, while LOB4DCG and LOBP4DCG (5) uses a subspace of maximum dimension $3n_{\text{want}} = 540$



of LOBP4DCG is closely related to the preconditioner used. We should note that the generic preconditioner (4.1) is not the natural preconditioner for TEST 1 to TEST 3. For plane wave-based calculations, it is more natural to use a proper scaled diagonal-like preconditioner proposed in [24]. In fact, K and M are not needed explicitly in plane wave-based calculations and they only exist in certain structural form, but K and M in our test were output from turboTDDFT runs. In such a case, the scaled diagonal preconditioner is lost once K and M are output explicitly as matrices. A future work is to compare them directly in turboTDDFT runs. Third, the advantage of BChebyDLR(ℓ) is that it does not need to apply preconditioners to solve linear equations. However, this convenience comes with a cost, namely that the current method is particularly effective only for the cases in which M is well-conditioned. Further research is required to make the method also efficient for M that has a relatively large condition number.

Example 4.2 When the K-matrix is indefinite, LOBP4DCG becomes not applicable. Therefore we can only report numerical results of Algorithm 2.2 without any comparison to LOBP4DCG. Since no other iterative methods seem to exploit the special structure of (1.2), known eigenvalue algorithms would treat (1.2) as a size $2n \times 2n$ standard eigenvalue problem, which is clearly an expensive thing to do. Therefore we do not compare with any general eigensolver here either. Instead, we use this example to demonstrate some common properties of BChebyDLR(ℓ), such as the effect of block sizes (see Fig. 2).

We apply BChebyDLR(ℓ) to the matrices listed as TEST 4, TEST 5, and TEST 6 in Table 1. Since the K-matrix for each of these tests contains negative eigenvalues, the first few eigenvalues of the simulated LREP of form (1.2) are purely imaginary. As shown in Fig. 2, BChebyDLR(ℓ) has no difficulty at all with the indefiniteness of K. This is also true when we vary the polynomial degree m, as shown in part of the next example (Fig. 3).

It is clear from Fig. 2 that the single vector version ($\ell=1$) is significantly slower than its blocked counterparts ($\ell>1$). However, when $n_{\rm want}$ is not large, as for the shown $n_{\rm want}=90$ case, using a too large block size may not lead to improved efficiency.

Example 4.3 We report the effect of the polynomial degree m in BChebyDLR(ℓ) on TEST 4 and TEST 6 in Fig. 3. Results on other tests are similar thus omitted. We let m vary from 8 to 38 with a stride length 3. All other parameters are fixed for these tests. Figure 3 plots the total CPU time in seconds, total number of iterations (#iter), and total number of matrix-vector products (#mvp) vs. the polynomial degree m.

As expected, the number of total iterations decreases as m increases, while the total number of matrix-vector products does not change monotonically with m. The figure shows that the total #mvp is not a deciding factor on total CPU time. When m is larger, the method likely uses more #mvp, but the significantly reduced #iter leads to smaller total CPU time. For these examples, the degree m around 25 to 35 all appear to be cost-effective; further increasing m over 40 usually does not have as significant effect as increasing m when it is below 15. The least efficient case is when m is too small. In comparison, the non-filtered case (m = 1), which in theory is



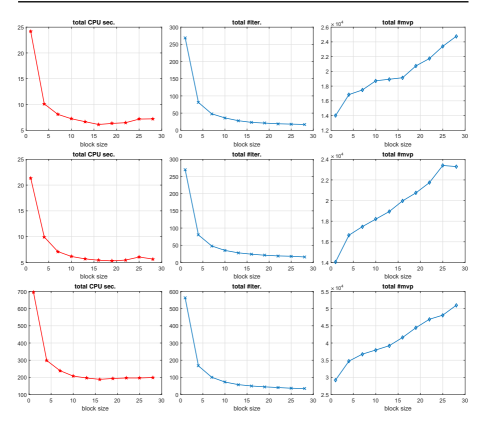


Fig. 2 Cost in computing the first $n_{\text{want}} = 90$ eigenpairs of TEST 4 (top), TEST 5 (middle), and TEST 6 (bottom). The filter degree is fixed at m = 25. The block size ℓ of BChebyDLR(ℓ) varies from 1 to 28 with a stride length 3

equivalent to the Lanczos method [18, 21, 22], is significantly slower than the filtered method with a moderate m. For the non-filtered method to be competitive in speed, one would need to use a much larger dimensional subspace (a number usually much greater than $2n_{\text{want}}$), but the memory constraint can make it impractical for larger problems.

As seen from Fig. 3, it is rather easy to choose and fix an m for the BChebyDLR method to be cost-effective, but we mention that [1] contains an interesting idea on adaptively updating m during the iterations.

Example 4.4 In the previous examples, we use the estimated β by Algorithm 3.2 as an upper bound for λ_n^2 . It works well in general. However, that β is not guaranteed to be a true upper bound. It may be smaller than λ_n^2 . Here we test the effectiveness of the adaptive strategy discussed at the end of Section 3. For this experiment, we



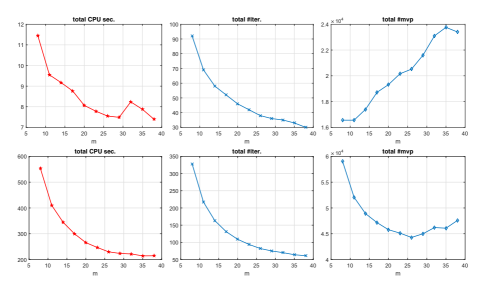


Fig. 3 Computing the first $n_{\text{want}} = 100$ eigenpairs of TEST 4 (top) and TEST 6 (bottom). The maximum subspace dimension used is 130. The filter degree m varies from 8 to 38 with a stride length 3. The block size $\ell = 10$ is used for all test runs

purposely begin with a β that underestimates λ_n^2 so that it can be updated by the adaptive strategy. We denote the resulting algorithm as a-BChebyDLR. Again we use the test cases TEST 1 to TEST 6. We compare a-BChebyDLR and BChebyDLR with block size $\ell=15$. The upper bound β and the parameters α and α_0 obtained in Algorithm 3.2 are used in BChebyDLR. We purposely choose an initial $\beta:=\beta_a=\lambda_{\max}(T_k)$, where T_k is symmetric tridiagonal matrix (3.5) computed in Algorithm 3.1, for a-BChebyDLR. This β is an underestimate, since $\lambda_{\max}(T_k) \leq \lambda_n^2$ owing to Eq. (3.7). Other parameters are fixed to be the same for both schemes. We plot the number of iterations and matrix-vector products in Fig. 4. This figure shows that even with an initial filter upper bound that underestimate the largest eigenvalue, one can readily make BChebyDLR work properly, as demonstrated by the result of a-BChebyDLR. The main trick is to simply update the filter upper bound to be the largest Ritz value ρ_{\max} (as defined in Eq. (3.7)), when it is found that $\rho_{\max} > \beta$.

An interesting feature of BChebyDLR is that the total iteration number #iter can be much smaller than the number of computed eigenpairs n_{want} , as seen in Table 2 and Figs. 2 and 3. This means that BChebyDLR, with suitable block size ℓ and degree m, can converge more than one eigenpair within one single iteration step. Similar effects of m and ℓ using Chebyshev filters for different applications from LREP were also observed in [31]. The high efficiency of Chebyshev filters, as constructed in [33, 34] and also here, has attracted attention and contributed to wide spread usage of Chebyshev filter based methods, e.g., in [4, 11, 13], for large-scale DFT calculations.



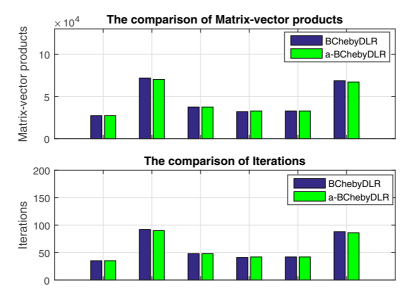


Fig. 4 Comparison of the number of iterations and the number of matrix-vector products by BChebyDLR and a-BChebyDLR (which employs the adaptive strategy), using the same block size $\ell=15$ and the same polynomial degree m=25 to compute the first $n_{\rm want}=180$ eigenpairs. The performances by both are comparable

5 Conclusion

We propose a Chebyshev-Davidson method BChebyDLR for solving the linear response eigenvalue problem (LREP). This method can effectively calculate the smallest positive eigenvalues and associated eigenvectors of LREP. In Theorem 2.2 we provide a convergence rate estimate for computing the smallest eigenvalue.

Numerical examples demonstrate that BChebyDLR is competitive to the recently proposed LOBP4DCG method [2]. BChebyDLR needs an upper bound that is no smaller than the largest λ_n^2 in order to be effective. Although the bound computed by Algorithm 3.2 usually works well, it needs conditions as specified in Theorem 3.1 to guarantee a strict upper bound. These conditions are non-trivial to verify by computation. We propose a simple adaptive strategy that can obtain an effective upper bound during the later filtering steps in BChebyDLR if the bound from Algorithm 3.2 is an underestimate of λ_n^2 for the first filtering step.

We emphasize that the proposed BChebyDLR works for both the definite (both K and M are positive definite) and indefinite (one of K and M is indefinite) cases (see Example 4.2), while in contrast LOBP4DCG is applicable only to the definite LREP.

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