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Numerical studies on structure-preserving algorithms for surface acoustic wave simulations

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ABSTRACT

We study the generalized eigenvalue problems (GEPs) derived from modeling the surface acoustic wave in piezoelectric materials with periodic inhomogeneity. The eigenvalues appear in the reciprocal pairs due to periodic boundary conditions in the modeling. By transforming the GEP into a T-palindromic quadratic eigenvalue problem (TPQEP), the reciprocal relationship of the eigenvalues can be maintained. In this paper, we outline four recently developed structure-preserving algorithms, SA, SDA, TSHIRA and GTSHIRA, for solving the TPQEP. Numerical comparisons on the accuracy and the computational costs of these algorithm are presented. The eigenvalues close to unit circle on the complex plane are of interest in the area of filter and sensor designs. Our numerical results show that the Arnoldi-type structure-preserving algorithms TSHIRA and GTSHIRA with "re-symplectic" and "re-bi-isotropic" processes, respectively, are as accurate as the SA and SDA algorithms, and more efficient in finding these eigenvalues.

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1. Introduction

In this paper we consider the generalized eigenvalue problem (GEP) of the form

$$\begin{bmatrix} M_1 & G\\ F^{\top} & 0 \end{bmatrix} \begin{bmatrix} \psi_i\\ \psi_\ell \end{bmatrix} + \lambda \begin{bmatrix} 0 & F\\ G^{\top} & M_2 \end{bmatrix} \begin{bmatrix} \psi_i\\ \psi_\ell \end{bmatrix} = 0,$$
(1)

where $M_1^{\top} = M_1 \in \mathbb{C}^{n \times n}, M_2^{\top} = M_2 \in \mathbb{C}^{m \times m}, F$ and $G \in \mathbb{C}^{n \times m}$ with $m \ll n$, and the subscript " \top " denotes the complex transpose. If M_1 and M_2 are nonsingular, then (1) can be reduced as the T-palindromic quadratic eigenvalue problem (TPQEP) of the form

$$\mathcal{P}(\lambda)x \equiv (\lambda^2 A_1^\top + \lambda A_0 + A_1)x = 0, \tag{2}$$

where

$$x = \psi_{\ell}, \qquad \psi_{i} = -M_{1}^{-1}(\lambda F + G)\psi_{\ell},$$

$$A_{1} = F^{\top}M_{1}^{-1}G, \qquad A_{0} = F^{\top}M_{1}^{-1}F + G^{\top}M_{1}^{-1}G - M_{2};$$
(3)

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or

$$x = \psi_i, \qquad \psi_\ell = -\lambda^{-1} M_2^{-1} (F^+ + \lambda G^+) \psi_i, A_1 = G M_2^{-1} F^\top, \qquad A_0 = F M_2^{-1} F^\top + G M_2^{-1} G^\top - M_1.$$
(4)

By taking the transpose of $\mathcal{P}(\lambda)$ in (2) and multiplying it by $1/\lambda^2$ it is easily seen that the eigenvalues of $\mathcal{P}(\lambda)$ appear in the reciprocal pairs $(\lambda, 1/\lambda)$ (including 0 and ∞). Since the nullity of $A_1 = GM_2^{-1}F^{\top}$ in (4) is larger or equal to n - m, $\mathcal{P}(\lambda)$ in (2) with A_0 and A_1 defined in (4) has n - m trivial zero and infinite eigenvalues which are not interested. We are only interested in finding $2m(\ll 2n)$ nontrivial eigenpairs of $\mathcal{P}(\lambda)$.

The GEP (1) can be solved by traditional methods such as QZ and Arnoldi method. But it does not guarantee that half of the computed eigenvalues lie inside of the unit circle and the others are outside [1]. For solving TPQEP (2) with small and dense matrices A_0 and A_1 , some pioneering works [2–4] have been done for preserving the reciprocity of the eigenvalues basing on a good linearization of (2) which transforms (2) into the form $\lambda Z^{\top} + Z$. Some structure-preserving methods [2,5,6] were proposed for solving ($\lambda Z^{\top} + Z$)u = 0. These methods require at least $\mathcal{O}(n^3)$ flops. Other structure-preserving algorithms can also be employed to solve (2). One method based on doubling algorithm was developed in [7] via the computation of a solvent of a nonlinear matrix equation associated with (2). Another structure-preserving algorithm based on ($\$ + \$^{-1}$)-transform [8] and Patel's approach [9] was developed in [10]. For problems with large and sparse matrices A_0 and A_1 , two structure-preserving Arnoldi-type algorithms, TSHIRA and GTSHIRA, were developed to search eigenvalues in a specified region of interests [10] where TSHIRA solves the standard \top -skew-Hamiltonian eigenvalue problem and GTSHIRA solves the generalized \top -skew-Hamiltonian eigenvalue problem. Both methods employed the ($\$ + \$^{-1}$)-transform and implicitly-restarted shift-and-invert Arnoldi method. However, in case eigenvectors are needed, an extra linear system has to be solved in TSHIRA.

The GEP (1) typically arises in many application areas including rail vibrations of fast train, surface acoustic wave (SAW) in filter design and crack modeling, etc., [11]. In these areas, an accurate and efficient eigensolver which preserves the reciprocal relationship of the associated eigenpairs is needed. In this paper, we would like to compare the accuracy and computational costs of the above mentioned algorithms for computing reciprocal eigenpairs in a SAW device [12]. The SAW filter plays an important role in telecommunication filters [13,14] and sensor technologies [15] etc. These filters are built on the physical property of piezoelectric materials, that electrical charges induce mechanical deformations and vice versa. The main component (or cell) of a SAW filter composes of a piezoelectric substrate and the input and output interdigital transducers (IDT). An input electrical signal from the input IDT produces a surface acoustic wave, traveling through periodically arranged electrodes and the output IDT picks up the output electrical signal. Depending on the material properties of the piezoelectric substrate (PZT) and the metallic electrode, and the gap length between the electrodes, frequencies in a desired range can be stopped or filtered out. In the filter design, it is important to know the stop band width and the center frequency f_c of the filter where $f_c = \frac{v_s}{\lambda_s}$ here v_s and λ_s are the wave velocity and wave length of the incident wave. The center frequency and stop band width can be determined by experiments or computation. In computational approach, the dispersion diagram needs to be generated in which a GEP of the form (1) associated with each frequency in the search range has to be solved [1].

This paper is organized as follows. We shall first introduce finite element modeling for a simple SAW resonator in Section 2. For more finite element simulations of piezoelectric devices in two dimension (2D) and three dimension (3D), one can refer to the works done by Allik, Koshiba, Lerch, Buchner and Mohamed etc., [16–19]. In Section 3, we introduce four structure-preserving algorithms developed in [7,10] to solve the TPQEP (2) and the GEP (1) resulted from our FEM model. Our numerical experiments in Section 4 compare the efficiency and accuracy of the structure-preserving algorithms for solving the GEP (1). Finally, we conclude the paper in Section 5.

2. Surface wave propagation

To model the wave propagation in a SAW device, we assume that a large number of electrodes are placed equally-spaced along a straight line on the PZT substrate. According to the Floquet–Bloch theory, one can reduce the problem to a single cell domain with one electrode by assuming the wave ψ is quasi-periodic of the form

$$\psi(x_1, x_2) = \psi_p(x_1, x_2)e^{(\alpha + i\beta)x_1}, \qquad \psi_p(x_1 + p, x_2) = \psi_p(x_1, x_2),$$

where x_1 is the wave propagation direction, p is the length of the unit cell (i.e. the periodic interval), α and β are the attenuation and phase shift along the wave propagation direction, respectively.

Let Ω denote the piezoelectric substrate with a single IDT as shown in Fig. 1, and Γ_{ℓ} and Γ_{r} denote the left and right boundary segments of Ω , respectively. For the general anisotropic PZT substrates, under the assumption of linear piezoelectric coupling, the elastic and electric fields interact following the general material constitution law below

$$T = c^{E}S - e^{\top}E,$$

$$D = eS + \varepsilon^{S}E,$$
(5)



Fig. 1. A 2D single cell domain of a LSAW resonator and boundary conditions.

where vectors *T*, *S*, *D* and *E* are the mechanical stress, strain, dielectric displacement and the electric field, respectively, and the matrices c^E , ε^S and *e* are the elasticity constant, dielectric constant and piezoelectric constant matrices measured at constant electric and constant strain fields at constant temperature. By applying the virtual work principle to the Eq. (5), the equilibrium state satisfies the following equation:

$$\int_{\Omega} (\delta S)^{\top} \left[c^{E} S + e^{\top} (\nabla \phi) \right] dV + \int_{\Omega} (\nabla \delta \phi)^{\top} \left[eS - \varepsilon^{S} (\nabla \phi) \right] dV + \int_{\Omega} (\delta u)^{\top} \rho \ddot{u} dV$$
$$= \int_{\Gamma_{\tilde{l}} \cup \Gamma_{r}} \left[(\delta u)^{\top} (T \cdot \vec{n}) + (\delta \phi)^{\top} (D \cdot \vec{n}) \right] dA, \tag{6}$$

where ρ is the mass density, $u = [u_1, u_2, u_3]^{\top}$ is the displacement vector, ϕ is the electric potential that satisfies $\nabla \phi = E, S = \begin{bmatrix} \frac{\partial u_1}{\partial x}, \frac{\partial u_2}{\partial y}, \frac{\partial u_3}{\partial z}, \frac{\partial u_2}{\partial z} + \frac{\partial u_3}{\partial y}, \frac{\partial u_3}{\partial x} + \frac{\partial u_1}{\partial z}, \frac{\partial u_1}{\partial y} + \frac{\partial u_2}{\partial x} \end{bmatrix}^{\top}$, and $\delta u, \delta \phi, \delta S$ are virtual displacement, potential and strain vectors, respectively. Let the notation $\psi = [u^{\top}, \phi]^{\top}$ and the subscript *i*, ℓ and *r* refer to nodal point index in the interior, the left boundary and the right boundary of the domain Ω , respectively. Using the periodic boundary conditions, proposed by Buchner [17],

$$T_r \cdot n_r = -\gamma T_\ell \cdot n_\ell, \qquad D_r \cdot n_r = -\gamma D_\ell \cdot n_\ell \quad \text{with } \gamma = e^{-(\alpha + i\beta)},$$

the finite element discretization to (6) on the domain Ω [1] can be written in the following matrix form

$$C(\omega)\psi \equiv [K - \omega^2 M + \iota\omega(\kappa_1 K + \kappa_2 M)]\psi = 0, \tag{7}$$

where $\kappa_1, \kappa_2 > 0$ are the viscous damping and mass damping respectively. By ordering the nodal unknown ψ according the order of subscripts ℓ , *i* and *r*, the matrices *K* and *M*, and the vector ψ can be partitioned as following:

$$K = \begin{bmatrix} K_{\ell\ell} & K_{i\ell}^{\top} & 0\\ K_{i\ell} & K_{ii} & K_{ir}\\ 0 & K_{ir}^{\top} & K_{rr} \end{bmatrix}, \qquad M = \begin{bmatrix} M_{\ell\ell} & M_{i\ell}^{\top} & 0\\ M_{i\ell} & M_{ii} & M_{ir}\\ 0 & M_{ir}^{\top} & M_{rr} \end{bmatrix}$$

where $K_{ii}, M_{ii} \in \mathbb{R}^{n \times n}, K_{\ell\ell}, K_{rr}, M_{\ell\ell}, M_{rr} \in \mathbb{R}^{m \times m}, K_{i\ell}, K_{ir}, M_{i\ell}, M_{ir} \in \mathbb{R}^{n \times m}$, and $\psi = [\psi_{\ell}^{\top}, \psi_{i}^{\top}, \psi_{r}^{\top}]^{\top}$ with $\psi_{i} \in \mathbb{C}^{n}, \psi_{\ell}, \psi_{r} \in \mathbb{C}^{m}(m \ll n)$. Obviously the matrix $C(\omega)$ in (7) can also be partitioned into

$$C(\omega) \equiv C \equiv \begin{bmatrix} C_{\ell\ell} & C_{i\ell}^{\top} & 0\\ C_{i\ell} & C_{ii} & C_{ir}\\ 0 & C_{ir}^{\top} & C_{rr} \end{bmatrix}.$$

By setting $\psi_r = \lambda \psi_\ell$, the Eq. (7) leads to the generalized eigenvalue problem

$$\begin{pmatrix} \begin{bmatrix} C_{ii} & C_{i\ell} \\ C_{ir}^{\top} & 0 \end{bmatrix} - \lambda \begin{bmatrix} 0 & C_{ir} \\ C_{i\ell}^{\top} & C_{bb} \end{bmatrix} \begin{pmatrix} \psi_i \\ \psi_\ell \end{bmatrix} = 0,$$

$$(8)$$

where $C_{bb} := C_{\ell\ell} + C_{rr}$.

Since the viscosity is small for PZT substrates and metals in SAW devices, the attenuation factor α of surface waves is close to zero. As a result, the propagation factor λ are generally near the unit circle thereafter denoted by \mathbb{U} . Furthermore, for frequency ω in the stopping band, the frequency shift parameter β shall be close to π when the periodic interval p (i.e. the domain width here) equals to half of the incident wave length λ_s . Therefore, we are interesting in finding λ close to \mathbb{U} , especially for those are near -1 on the complex plane. Notice that eigenvalues of (2) appear in the reciprocal pairs

 $(\lambda, 1/\lambda)$. In the following sections, we aim to discuss the efficiency and accuracy of the structure-preserving algorithms [7,10] for solving the eigencurves $\lambda(\omega)$ and the associated eigenvectors of (8).

3. Structure-preserving algorithms

In this section, we shall introduce four structure-preserving algorithms developed in [7,10] to solve the TPOEP (2) and discuss the computation costs of these algorithms in solving the GEP (1). In the following, we suppose m reciprocal pairs of eigenvalues near \mathbb{U} are desired.

3.1. Structure-preserving doubling algorithm

For solving the TPQEP (2) with $A_0, A_1 \in \mathbb{C}^{m \times m}$ defined in (3), a structure-preserving doubling algorithm (SDA) was developed in [7] via the computation of a solvent of a nonlinear matrix equation associated with (2). That is $\mathcal{P}(\lambda)$ can be factorized as

$$\mathcal{P}(\lambda) = (\lambda A_1^{\top} - X)X^{-1}(\lambda X - A_1)$$
(9)

for some nonsingular X with $X^{\top} = X$ if and only if X satisfies the following nonlinear matrix equation (NME):

$$A_1^{\dagger} X^{-1} A_1 + X + A_0 = 0.$$

Combining SDA in [7], the GEP (1) can be solved by Algorithm 1. The advantages of Algorithm 1 are as following: (i) the computed eigenvalues are guaranteed to appear in reciprocal pair since the eigenvalues of the matrix pencils $\lambda A_1^{\top} - X$ and $\lambda X - A_1$, which are reciprocal pairs, are the eigenvalues of $\mathcal{P}(\lambda)$ in (9) and (ii) the convergence rate of the SDA is proved to be guadratic [7] if there are no eigenvalues of $\mathcal{P}(\lambda)$ located on unit circle.

Algorithm 1 GE_SDA

Input: matrices *F*, *G*, M_2 and M_1 , tolerance η and the number m of desired eigenvalues.

Output: eigenpairs
$$\{(\gamma_j, [(\psi_{\ell,j}^{(1)})^{\top}, (\psi_{\ell,j}^{(1)})^{\top}]^{\top}), (\gamma_j^{-1}, [(\psi_{\ell,j}^{(2)})^{\top}, (\psi_{\ell,j}^{(2)})^{\top}]^{\top})\}_{j=1}^{\mathfrak{m}}$$
 of (1).

1: Compute
$$A_0 = F^+ M_1^{-1}F + G^+ M_1^{-1}G - M_2$$
 and $A_1 = F^+ M_1^{-1}G$.

2: Set
$$k = 0$$
, $Y_k = A_1$, $X_k = -A_0$ and $Z_k = 0$.

4: Compute
$$Y_{k+1} = Y_k(X_k - Z_k)^{-1}Y_k$$
, $X_{k+1} = X_k - Y_k^{\top}(X_k - Z_k)^{-1}Y_k$, and $Z_{k+1} = Z_k + Y_k(X_k - Z_k)^{-1}Y_k^{\top}$;

5: Set
$$k = k + 1$$
;

6: **until**
$$||X_k - X_{k-1}|| \le \eta ||X_k||$$

7: Compute the left and right eigenpairs $\{(\lambda_j, \psi_{\ell,j}^{(1)}), (\lambda_j, \psi_{\ell,j}^{(r)})\}_{j=1}^m$ of $X_k \psi_\ell = \lambda A_1^\top \psi_\ell$;

8: Choose the eigenpairs which associated eigenvalues are near the unit circle, said $\{(\lambda_j, \psi_{\ell,j}^{(1)}), (\lambda_j, \psi_{\ell,j}^{(r)})\}_{j=1}^{\mathfrak{m}}$; 9: Solve $(\lambda_j X_k - A_1)\psi_{\ell,j}^{(2)} = X_k \psi_{\ell,j}^{(r)}$ and set $\gamma_j = \lambda_j^{-1}$ for $j = 1, ..., \mathfrak{m}$;

10: Compute

$$\psi_{i,j}^{(1)} = -M_1^{-1} \left(\gamma_j F \psi_{\ell,j}^{(1)} + G \psi_{\ell,j}^{(1)} \right), \ \psi_{i,j}^{(2)} = -M_1^{-1} \left(\gamma_j^{-1} F \psi_{\ell,j}^{(2)} + G \psi_{\ell,j}^{(2)} \right)$$

for $j = 1, \ldots, m$.

Next, let us discuss the computational costs of Algorithm 1. To mimic the computation cost in the LU factorization of the matrix M_1 obtained from finite element discretization, we reorder the nodal indices so that the matrix M_1 has narrower band structure. Let $M_1 = LU$ be the LU factorization of M_1 . Then, computing A_0 and A_1 in Step 7 of Algorithm 1 requires solving $\widetilde{F} \equiv U^{-1}L^{-1}F$ and $\widetilde{G} \equiv U^{-1}L^{-1}G$, and matrix multiplications of $F^{\top}\widetilde{F}$, $G^{\top}\widetilde{G}$ and $F^{\top}\widetilde{G}$. In Steps 3–6, one LU factorization $(2m^3/3)$ flops), two forward and back substitutions (4m³ flops) and three matrices multiplications (6m³ flops) are required for each iterate k. Next, computing the left and right eigenpairs in Step 7 and solving $\psi_{\ell,j}^{(2)}$ in Step 9 take 100m³ flops and 2mm³/3 flops, respectively. Finally, it also requires $2\mathfrak{m}$ forward and back substitutions to compute $\{\psi_{i,j}^{(1)}, \psi_{i,j}^{(2)}\}_{j=1}^{\mathfrak{m}}$ in Step 10. The total cost of Algorithm is summarized in Table 1.

3.2. Structure-preserving algorithm

Another structure-preserving algorithm (SA) developed in [10] is based on the $(\delta + \delta^{-1})$ -transform [8] and Patel's approach [9] for solving the TPQEP (2) with $A_0, A_1 \in \mathbb{C}^{m \times m}$ defined in (3). The idea is, first, to linearize the TPQEP as the following special GEP:

$$(\mathcal{M} - \lambda \mathcal{L}) \begin{bmatrix} x \\ y \end{bmatrix} = 0, \tag{10}$$

Table 1

The computational costs of GE_SDA and GE_SA where k denotes the total iterations to obtain convergent X_k in Lines 3.1–3.1 of GE_SDA.

| Compute $M_1 = III$ | | GE SDA | GE SA |
|--------------------------------------|-------------------------|---|------------------------|
| eompare mi 20 | | | |
| | | 1 | 1 |
| Compute A_0, A_1 | Solve $Lx = b_1$ | 2 <i>m</i> | 2 <i>m</i> |
| | Solve $Ux = b_2$ | 2 <i>m</i> | 2 <i>m</i> |
| | Compute $F^{\top}d_1$ | 2 <i>m</i> | 2 <i>m</i> |
| | Compute $G^{\top}d_2$ | т | т |
| Compute $\psi_i^{(1)}, \psi_i^{(2)}$ | Solve $Lx = b_1$ | 2m | 2m |
| | Solve $Ux = b_2$ | 2m | 2m |
| | Compute Fe ₁ | 2m | 2m |
| | Compute Ge ₂ | 2m | 2m |
| Solve dense TPQEP | | $(100 + \frac{32}{3}k + \frac{2}{3}m)m^3$ flops | 50m ³ flops |

where $\lambda y = A_1 x$, and

$$\mathcal{M} = \begin{bmatrix} A_1 & 0\\ -A_0 & -I \end{bmatrix}, \qquad \mathcal{L} = \begin{bmatrix} 0 & I\\ A_1^\top & 0 \end{bmatrix}.$$
(11)

Obviously, the matrix pencil $\mathcal{M} - \lambda \mathcal{L}$ is \top -symplectic, i.e., it satisfies $\mathcal{M}\mathcal{J}\mathcal{M}^{\top} = \mathcal{L}\mathcal{J}\mathcal{L}^{\top}$ where $\mathcal{J} = \begin{bmatrix} 0 & l_m \\ -l_m & 0 \end{bmatrix}$. As a result, the eigenvalues of $(\mathcal{M}, \mathcal{L})$ appear in the reciprocal pairs $(\lambda, 1/\lambda)$. Secondly, the $(\mathcal{S} + \mathcal{S}^{-1})$ -transform is applied on $\mathcal{M} - \lambda \mathcal{L}$ and the pencil is now transformed into a \top -skew-Hamiltonian pencil $\mathcal{K} - \mu \mathcal{N}$, i.e., $(\mathcal{K}\mathcal{J})^{\top} = -\mathcal{K}\mathcal{J}, (\mathcal{N}\mathcal{J})^{\top} = -\mathcal{N}\mathcal{J}$:

$$\mathcal{K} - \mu \mathcal{N} \equiv \left[(\mathcal{L} \mathcal{J} \mathcal{M}^{\top} + \mathcal{M} \mathcal{J} \mathcal{L}^{\top}) - \mu \mathcal{L} \mathcal{J} \mathcal{L}^{\top} \right] \mathcal{J}^{\top} = \begin{bmatrix} A_0 & A_1^{\top} - A_1 \\ A_1 - A_1^{\top} & A_0 \end{bmatrix} - \mu \begin{bmatrix} -A_1 & 0 \\ 0 & -A_1^{\top} \end{bmatrix}.$$
(12)

The two eigenvalues λ and μ are then related by the relationship $\mu = \lambda + 1/\lambda$. The relationship between eigenpairs of the TPQEP in (2) and the \top -skew-Hamiltonian pair (\mathcal{K} , \mathcal{N}) in (12) is stated in the following theorem.

Theorem 3.1 ([10]). Let $(\mathcal{K}, \mathcal{N})$ be defined in (12). If $z_s = [z_1^{\top}, z_2^{\top}]^{\top}$ with $z_1, z_2 \in \mathbb{C}^m$ is an eigenvector of $(\mathcal{K}, \mathcal{N})$ corresponding to eigenvalue μ and ν satisfies $\nu + \frac{1}{\nu} = \mu$, then $\frac{1}{\nu}z_1 - z_2$ and $\nu z_1 - z_2$ are eigenvectors of the TPQEP in (2) corresponding to eigenvalues ν and $\frac{1}{\nu}$, respectively.

Finally, based on Patel's approach [9], the matrix pair $(\mathcal{K}, \mathcal{N})$ can further be reduced to a block triangular structure as following

$$\mathcal{K} := Q^{\top} \mathcal{K} Z = \begin{bmatrix} K_{11} & K_{12} \\ 0 & K_{11}^{\top} \end{bmatrix}, \qquad \mathcal{N} := Q^{\top} \mathcal{N} Z = \begin{bmatrix} N_{11} & N_{12} \\ 0 & N_{11}^{\top} \end{bmatrix}, \tag{13}$$

where $K_{11} \in \mathbb{C}^{m \times m}$ is upper Hessenberg, $N_{11} \in \mathbb{C}^{m \times m}$ is upper triangular, and Q, Z are unitary satisfying

$$Q = \mathcal{J}^{\top} Z \mathcal{J}.$$

We then apply the QZ algorithm to (K_{11}, N_{11}) for computing the *m* eigenpairs $\{(\mu_k, y_k)\}_{k=1}^m$. Consequently, $\{(\mu_k, Z \begin{bmatrix} y_k \\ 0 \end{bmatrix})\}_{k=1}^m$ are the *m* eigenpairs of $(\mathcal{K}, \mathcal{N})$. Combining the above procedures and the structure-preserving algorithm in [10], the GEP (1) can be solved by Algorithm 2.

The computational costs in Steps 1 and 9 of Algorithm 2 are the same that in Steps 1 and 10 of Algorithm 1. The SA processes in Steps 2–8 of Algorithm 2 require approximately $50m^3$ flops [10] to compute the eigenpairs of the TPQEP (2) with small size matrices A_0 and A_1 in (3). The comparison of the computation costs for GE_SDA and GE_SA is listed in Table 1.

3.3. *⊤*-skew-Hamiltonian implicit-restarted Arnoldi algorithm

In the above mentioned GE_SDA and GE_SA algorithms, the GEP (1) is transformed into the TPQEP (2) through equations in (3) where $M_1^{-1}F$ and $M_1^{-1}G$ are solved by LU factorization on the matrix M_1 . The computation costs in this step increase in the amount of 2m times n^2 . Since the GE_SDA and GE_SA algorithms are then working on the TPQEP where the size of matrices is $m \times m$, $m \ll n$, the computation cost in solving the TPQEP is relatively small.

In the following, we introduce two Arnoldi-type algorithms in which the GEP (1) is transformed into the TPQEP (2) through equations in (4). Since the matrix size *m* of M_2 is much smaller, the cost in solving $M_2^{-1}F^{\top}$ and $M_2^{-1}G^{\top}$ by LU

Algorithm 2 GE_SA

- **Input:** matrices *F*, *G*, *M*₂ and *M*₁, and the number m of desired eigenvalues. **Output:** eigenpairs { $(\gamma_j, [(\psi_{i,j}^{(1)})^\top, (\psi_{l,j}^{(1)})^\top]^\top), (\gamma_j^{-1}, [(\psi_{i,j}^{(2)})^\top, (\psi_{l,j}^{(2)})^\top]^\top)$ }_{j=1} of (1). 1: Compute $A_0 = F^\top M_1^{-1}F + G^\top M_1^{-1}G M_2$ and $A_1 = F^\top M_1^{-1}G$.
- 2: Form the pair $(\mathcal{K}, \mathcal{N})$ as in (12);
- 3: Reduce $(\mathcal{K}, \mathcal{N})$ to block upper triangular forms in (13) using unitary transformations;
- 4: Compute eigenpairs $\{(\mu_k, y_k)\}_{k=1}^m$ of (K_{11}, N_{11}) defined in (13) by using the QZ algorithm; 5: Compute eigenvalues v_k and v_k^{-1} of $\mathcal{P}(\lambda)$ by solving $v^2 \mu_k v + 1 = 0$;
- 6: Choose the eigenvalues which are near the unit circle, said $\{v_{k_i}, v_{k_i}^{-1}\}_{i=1}^{m}$;
- 7: Compute $z_j = Z \begin{bmatrix} y_{k_j} \\ 0 \end{bmatrix} \equiv \begin{bmatrix} z_{j1} \\ z_{j2} \end{bmatrix}, j = 1, 2, \dots, \mathfrak{m};$
- 8: Compute eigenvectors $\psi_{\ell,j}^{(1)} \equiv \gamma_j^{-1} z_{j1} z_{j2}$ and $\psi_{\ell,j}^{(2)} \equiv \gamma_j z_{j1} z_{j2}$ corresponding to eigenvalues $\gamma_j \equiv v_{k_j}$ and $v_{k_j}^{-1}$, respectively, for $j = 1, 2, \ldots, m$;
- 9: Compute

$$\psi_{i,j}^{(1)} = -M_1^{-1} \left(\gamma_j F \psi_{\ell,j}^{(1)} + G \psi_{\ell,j}^{(1)} \right), \ \psi_{i,j}^{(2)} = -M_1^{-1} \left(\gamma_j^{-1} F \psi_{\ell,j}^{(2)} + G \psi_{\ell,j}^{(2)} \right)$$

for $j = 1, \ldots, \mathfrak{m}$.

factorization of M_2 can now be ignored. Following the same idea in Section 3.2, the TPQEP (2) with $A_0, A_1 \in \mathbb{C}^{n \times n}$ is also transformed into the \top -skew-Hamiltonian pencil $\mathcal{K} - \mu \mathcal{N}$ through the Eqs. (10) and (12) with $\mathcal{J} = \begin{bmatrix} 0 & I_n \\ -I_n & 0 \end{bmatrix}$. Instead of taking Patel's approach, we seek the eigenvalues of the matrix pair $(\mathcal{K}, \mathcal{N})$ by some implicit-restart Arnoldi algorithms. Although the Arnoldi algorithm is working on the matrices with size $2n \times 2n$ now, saving on computation costs is expected when fast convergence of the Arnoldi iterations can be achieved. In the following, we sketch the key steps and theorems that are employed in developing Arnoldi algorithm.

Let τ be a shift value and $\tau \notin \sigma(\mathcal{M}, \mathcal{L})$ where $\sigma(\mathcal{A}, \mathcal{B})$ denotes the set of all eigenvalues of any matrix pair $(\mathcal{A}, \mathcal{B})$. Then, we have $\mu_0 \equiv \tau + \tau^{-1} \notin \sigma(\mathcal{K}, \mathcal{N})$. Define the shift-invert transformation $\widehat{\mathcal{K}} - \widehat{\mu}\widehat{\mathcal{N}}$ for $\mathcal{K} - \mu\mathcal{N}$ with $\widehat{\mu} = \frac{1}{\mu - \mu_0}$ and

$$\widehat{\mathcal{K}} \equiv -\tau \,\mathcal{N} = \tau \begin{bmatrix} A_1^\top & 0\\ 0 & A_1 \end{bmatrix},\tag{14a}$$

$$\widehat{\mathcal{N}} \equiv -\tau(\mathcal{K} - \mu_0 \mathcal{N}) = (\mathcal{M} - \tau \mathcal{L}) \mathcal{J} \left(\mathcal{M}^\top - \tau \mathcal{L}^\top \right) \mathcal{J}^\top, \tag{14b}$$

where $\widehat{\mathcal{K}}$ and $\widehat{\mathcal{N}}$ are \top -skew-Hamiltonian. Furthermore, from the definition of $\widehat{\mathcal{N}}$ in (14b), $\widehat{\mathcal{N}}$ can be factorized as $\widehat{\mathcal{N}}$ = $\mathcal{N}_1 \mathcal{N}_2$, where

$$\mathcal{N}_1 = \mathcal{M} - \tau \mathcal{L}, \qquad \mathcal{N}_2 = \mathcal{J}(\mathcal{M}^\top - \tau \mathcal{L}^\top) \mathcal{J}^\top$$
 (15)

are nonsingular and satisfy $\mathcal{N}_2^\top \mathcal{J} = \mathcal{J} \mathcal{N}_1$. The GEP $\widehat{\mathcal{K}} z = \widehat{\mu} \widehat{\mathcal{N}} z$ is then equivalent to the eigenvalue problem $\mathscr{B} \tilde{z} = \widehat{\mu} \tilde{z}$, where

$$\mathscr{B} \equiv \mathscr{N}_1^{-1} \widehat{\mathscr{K}} \mathscr{N}_2^{-1} \tag{16}$$

is \top -skew-Hamiltonian, i.e., $\mathcal{JB}^{\top} = \mathcal{B}\mathcal{J}$, and $\tilde{z} = \mathcal{N}_2 \hat{z}$. Now, according to the following two main theorems proved in [10,20], the T-skew-Hamiltonian implicitly-restarted Arnoldi (TSHIRA) algorithm as shown in Algorithm 4 can be employed to solve this eigenvalue problem.

Let us define the Krylov matrix with respect to u_1 by

 $K_i \equiv K_i[\mathcal{B}, u_1] = [u_1, \mathcal{B}u_1, \ldots, \mathcal{B}^{j-1}u_1], \quad 1 \leq j \leq n.$

The two main theorems in [10,20] are as follows:

Theorem 3.2 ([20]). Let $\mathcal{B} \in \mathbb{C}^{2n \times 2n}$ be \top -skew-Hamiltonian and K_i be a Krylov matrix with rank $(K_i) = j$. Then span (K_i) is \top -isotropic and if $K_i = U_i R_i$ is a QR-factorization, then

$$\mathcal{B}U_{i} = U_{i}H_{i} + \tilde{u}_{i+1}e_{i}^{\top},$$

where $H_j \in \mathbb{C}^{j \times j}$ is unreduced upper Hessenberg, $U_j \in \mathbb{C}^{2n \times j}$ is orthonormal and \top -isotropic, and $\tilde{u}_{j+1} \in \mathbb{C}^{2n}$ is a suitable vector such that

 $U_i^H \tilde{u}_{j+1} = 0$ and $U_i^\top \mathcal{J} \tilde{u}_{j+1} = 0.$

Theorem 3.3 ([10,20]). Let $\mathcal{B} \in \mathbb{C}^{2n \times 2n}$ be \top -skew-Hamiltonian. If rank(K_n) = n, then there is a unitary \top -symplectic matrix \mathcal{U} with $\mathcal{U}e_1 = u_1$ such that

$$\mathcal{U}^{H}\mathcal{B}\mathcal{U} = \begin{bmatrix} H_{n} & S_{n} \\ 0 & H_{n}^{\top} \end{bmatrix},$$

where $H_n \equiv [h_{ij}]$ is unreduced upper Hessenberg and S_n is \top -skew-symmetric.

Based on Theorem 3.2, the jth step of TSHIRA is given by

$$h_{j+1,j}u_{j+1} = \mathcal{B}u_j - \sum_{i=1}^{J} h_{ij}u_i,$$
(17)

where $h_{ij} = u_i^H \mathcal{B}u_j$, i = 1, ..., j and $h_{j+1,j} > 0$ is chosen so that $||u_{j+1}||_2 = 1$. In order to guarantee the \top -isotropic property of the space span $\{u_1, ..., u_{j+1}\}$ is preserved within machine precision, reorthogonalizing u_{j+1} against $\mathcal{J}U_j$ is necessary. As a result, the Eq. (17) is modified into

$$h_{j+1,j}u_{j+1} = \mathscr{B}u_j - \sum_{i=1}^j h_{ij}u_i - \sum_{i=1}^j t_{ij}\mathscr{G}\bar{u}_i,$$

where $t_{ij} = -u_i^{\top} \mathcal{JB}u_j$, i = 1, ..., j. The above procedure is stated in Algorithm 3.

Finally, we present TSHIRA with Krylov–Schur restart to solve the eigenvalue problem $\mathscr{B}\widetilde{z} = \widehat{\mu}\widetilde{z}$ in Algorithm 4. Once the eigenpair $(\widehat{\mu}, \widetilde{z})$ is obtained, one can recover the eigenpair (μ, z) of $(\mathscr{K}, \mathscr{N})$ from the relationship $\widehat{u} = \frac{1}{\mu - \mu_0}$ and the solution of the linear system $\mathscr{N}_2 z = \widetilde{z}$. The reciprocal eigenpair $(\lambda, \frac{1}{\lambda})$ and the associated eigenvectors of the TPQEP (2) are then followed from Theorem 3.1.

Algorithm 3 The *j*th ⊤-isotropic Arnoldi step

Input: ⊤-skew-Hamiltonian *B* and $U_j = [u_1, \dots, u_j]$ with $U_j^H U_j = I_j$ and $U_j^T \mathscr{G} U_j = 0$. **Output:** $[h_{1,j}, \dots, h_{j+1,j}]$ and u_{j+1} . 1: Compute $u_{j+1} = \mathscr{B} u_j$; 2: **for** $i = 1, \dots, j$ **do** 3: $h_{ij} = u_i^H u_{j+1}, u_{j+1} = u_{j+1} - h_{ij} u_i$ 4: **end for** 5: **for** $i = 1, \dots, j$ **do** 6: $t_{ij} = u_i^T \mathscr{G}^T u_{j+1}, u_{j+1} = u_{j+1} - t_{ij} \mathscr{G} \bar{u}_i$ 7: **end for** 8: Set $h_{j+1,j} := ||u_{j+1}||_2$ and $u_{j+1} := u_{j+1}/h_{j+1,j}$.

Algorithm 4 [8] TSHIRA for solving $\mathcal{B}\tilde{z} = \widehat{\mu}\tilde{z}$

Input: \top -skew-Hamiltonian matrix \mathcal{B} with starting vector u_1 .

Output: eigenpairs $(\widehat{\mu}_i, \widetilde{z}_i)$, $i = 1, ..., \mathfrak{m}$ of \mathcal{B} .

1: Use Algorithm 3 with starting vector u_1 to generate the mth step of \top -isotropic Arnoldi decomposition: $\mathcal{B}U_m = U_m H_m + h_{m+1,m} u_{m+1} e_m^\top$;

2: repeat

3: Use Algorithm 3 to extend the mth step of \top -isotropic Arnoldi decomposition to the $(\mathfrak{m} + p)$ th step of \top -isotropic Arnoldi factorization:

 $\mathcal{B}U_{\mathfrak{m}+p} = U_{\mathfrak{m}+p}H_{\mathfrak{m}+p} + h_{\mathfrak{m}+p+1,\mathfrak{m}+p}u_{\mathfrak{m}+p+1}e_{\mathfrak{m}+p}^{\top}.$

4: Use Krylov–Schur restarting scheme [21,22] to reform a new ⊤-isotropic Arnoldi decomposition with order m.

5: **until** wanted m eigenpairs of *B* are convergent

3.4. Generalized *¬*-skew-Hamiltonian implicitly-restarted Arnoldi algorithm

Recall that an additional linear system $N_2 z = \tilde{z}$ has to be solved for recovering the eigenpair (μ, z) of $(\mathcal{K}, \mathcal{N})$ when TSHIRA is employed to solve the GEP $\hat{\mathcal{K}} z = \hat{\mu} \hat{\mathcal{N}} z$ in (14). This may result in losing some accuracy of the eigenvector z. In order to eliminate this extra computational cost and to prevent the inaccuracy, a generalized \top -skew-Hamiltonian implicitly-restarted Arnoldi (GTSHIRA) algorithm is proposed in [10]. The idea is to solve the GEP $\hat{\mathcal{K}} z = \hat{\mu} \hat{\mathcal{N}} z$ in (14) directly through bi-reorthogonalization and bi- \top -isotropic processes. The GTSHIRA algorithm is based on following two theorems.

Theorem 3.4 ([10]). Let $\mathcal{B} \equiv \mathcal{N}_1^{-1} \widehat{\mathcal{K}} \mathcal{N}_2^{-1}$ with $\widehat{\mathcal{N}} = \mathcal{N}_1 \mathcal{N}_2$ be \top -skew-Hamiltonian. Let $K_j \equiv K_j[\mathcal{B}, u_1]$ be the Krylov matrix with rank $(K_j) = j$. If

$$\mathcal{N}_2^{-1}K_j = Z_j R_{2,j}$$
 and $\mathcal{N}_1 K_j = Y_j R_{1,j}$

are QR-factorizations, where Z_j , $Y_j \in \mathbb{C}^{2n \times j}$ are orthonormal and $R_{2,j}$, $R_{1,j}$ are nonsingular upper triangular, then we have

$$\widehat{\mathcal{K}}Z_j = Y_j\widehat{H}_j + \widehat{y}_{j+1}e_j^\top$$
(18)

and

$$\widehat{\mathcal{N}}Z_j = Y_j\widehat{R}_j,\tag{19}$$

where $\widehat{H}_j \in \mathbb{C}^{j \times j}$ is unreduced upper Hessenberg, $\widehat{R}_j \in \mathbb{C}^{j \times j}$ is nonsingular upper triangular, and Y_j and Z_j are \top -bi-isotropic such that

$$Y_j^H \widehat{y}_{j+1} = 0$$
 and $Z_j^\top \mathscr{J} \widehat{y}_{j+1} = 0$,

for a suitable $\widehat{y}_{i+1} \in \mathbb{C}^{2n}$.

Theorem 3.5 ([10]). Let $\mathcal{B} = \mathcal{N}_1^{-1} \widehat{\mathcal{K}} \mathcal{N}_2^{-1}$ with $\widehat{\mathcal{N}} = \mathcal{N}_1 \mathcal{N}_2$ be \top -skew-Hamiltonian and $K_n \equiv K_n[\mathcal{B}, u_1]$ be the Krylov matrix with rank $(K_n) = n$. Then there are unitary matrices \mathcal{U} and \mathcal{V} satisfying $\mathcal{V} = \mathcal{J}^\top \mathcal{U} \mathcal{J}$, $\mathcal{U} e_1 = u_1$ and $\mathcal{V} e_1 = \mathcal{N}_1 u_1 / || \mathcal{N}_1 u_1 ||_2$ such that

$$\boldsymbol{\mathcal{V}}^{\top}\widehat{\boldsymbol{\mathcal{K}}}\boldsymbol{\mathcal{U}} = \begin{bmatrix} \widehat{H}_n & \widehat{S}_n \\ \mathbf{0} & \widehat{H}_n^{\top} \end{bmatrix}, \qquad \boldsymbol{\mathcal{V}}^{\top}\widehat{\boldsymbol{\mathcal{N}}}\boldsymbol{\mathcal{U}} = \begin{bmatrix} \widehat{R}_n & \widehat{T}_n \\ \mathbf{0} & \widehat{R}_n^{\top} \end{bmatrix}$$

where \widehat{H}_n is unreduced upper Hessenberg, \widehat{R}_n is nonsingular upper triangular and \widehat{S}_n , \widehat{T}_n are \top -skew-symmetric.

Based on Theorem 3.4 and assuming that the first (j - 1)th step in GTSHIRA follows the generalized \top -isotropic Arnoldi process, i.e.,

$$\widehat{\mathcal{N}}Z_{j-1} = Y_{j-1}\widehat{R}_{j-1},\tag{20}$$

by comparing the *j*th columns of both sides in (19) at the *j*th step, one has

$$\widehat{\mathcal{N}}z_j = \sum_{i=1}^{j-1} \widehat{r}_{ij} y_i + \widehat{r}_{jj} y_j.$$
(21)

With (20), (21) can be rewritten as

$$\widehat{r_{jj}}^{-1}z_j = \widehat{\mathcal{N}}^{-1}y_j - \sum_{i=1}^{j-1}\widetilde{r_{ij}}z_i,$$
(22)

where

$$[\widetilde{r}_{1j},\ldots,\widetilde{r}_{j-1,j}]^{\top}:=-\widehat{r}_{jj}^{-1}\widehat{R}_{j-1}^{-1}[\widehat{r}_{1j},\ldots,\widehat{r}_{j-1,j}]^{\top},$$

and \hat{r}_{ij} in (22) is chosen so that $||z_j||_2 = 1$. Since $Z_i^H Z_j = I_j$, the coefficient \tilde{r}_{ij} in (22) can be evaluated by

$$\widetilde{r}_{ij} = z_i^H \widehat{\mathcal{N}}^{-1} y_j, \quad i = 1, \dots, j-1.$$

Finally, from (18), the vector y_{j+1} in the *j*th step of the generalized \top -isotropic Arnoldi process is given by

$$\widehat{h}_{j+1,j}y_{j+1} = \widehat{\mathcal{K}}z_j - \sum_{i=1}^j \widehat{h}_{ij}y_i,$$

where

$$\widehat{h}_{ij} = y_i^H \widehat{\mathcal{K}} z_j,$$

and $\widehat{h}_{j+1,j} > 0$ is chosen so that $||y_{j+1}||_2 = 1$.

Notice that, in theory, z_j and y_{j+1} are orthogonal to $\mathcal{J}\bar{Y}_j$ and $\mathcal{J}\bar{Z}_j$, respectively, in exact arithmetic. However, in practice, roundoff errors may cause $y_i^{\top} \mathcal{J}^{\top} z_j$ and $z_i^{\top} \mathcal{J}^{\top} y_{j+1}$, i = 1, ..., j, to be some nonzero tiny values. Therefore, in order to preserve the \top -bi-isotropic property of Y_j and Z_j , reorthogonalization of z_j against $\mathcal{J}\bar{Y}_j$ or y_{j+1} against $\mathcal{J}\bar{Z}_j$ is needed. Summarizing above processes, we state the *j*th step of the generalized \top -isotropic Arnoldi process in Algorithm 5. The reorthogonalization

Algorithm 5 [8] The *j*th generalized ⊤-isotropic Arnoldi step

Input: \top -skew-Hamiltonian $\widehat{\mathcal{K}}$ and $\widehat{\mathcal{N}}$, upper triangular R(1:j-1,1:j-1), $Y_i = [y_1,\ldots,y_i]$ and $Z_{i-1} = [z_1,\ldots,z_{i-1}]$ with $Y_{j}^{H}Y_{j} = I_{j}, Z_{j-1}^{H}Z_{j-1} = I_{j-1}$ and $Y_{j}^{\top} \mathcal{J}Z_{j-1} = 0$. **Output:** $[h_{1,j}, \ldots, h_{j+1,j}], R(1:j,j), y_{j+1} \text{ and } z_j.$ 1: Solve $\widehat{\mathcal{N}} z_i = y_i$; 2: **for** $i = 1, \ldots, j - 1$ **do** 3: $\widehat{r}_{ij} = z_i^H z_j, \ z_j = z_j - \widehat{r}_{ij} z_i$ 4: end for 5: Reorthogonalize z_i to $\Im \overline{Y}_i$ as following for-loop in Steps 6–8: 6: **for** i = 1, ..., j **do** $s_{ij} = y_i^{\top} \mathcal{J}^{\top} z_j, \ z_j = z_j - s_{ij} \mathcal{J} \bar{y}_i$ 7: 8: end for 9: Set $R(j, j) := ||z_j||_2^{-1}, z_j := R(j, j)z_j$ and $R(1:j-1,j) := -R(j,j)R(1:j-1,1:j-1)[\widehat{r}_{1i},\ldots,\widehat{r}_{i-1,j}]^{\top}$ 10: Compute $y_{i+1} = \mathcal{K}z_i$; 11: **for** i = 1, ..., j **do** 12: $h_{ij} = y_i^H y_{j+1}, y_{j+1} = y_{j+1} - h_{ij} y_i$ 13: end for 14: Reorthogonalize y_{j+1} to $\mathcal{J}\overline{Z}_j$ as following for-loop in Steps 15–17: 15: **for** i = 1, ..., j **do** 16: $t_{ij} = z_i^{\top} \mathcal{J}^{\top} y_{j+1}, \ y_{j+1} = y_{j+1} - t_{ij} \mathcal{J} \bar{z}_i$ 17: end for 18: Set $h_{j+1,j} := \|y_{j+1}\|_2$ and $y_{j+1} := y_{j+1}/h_{j+1,j}$.

Algorithm 6 [8] GTSHIRA for solving $\widehat{\mathcal{K}}z = \widehat{\mu}\widehat{\mathcal{N}}z$

Input: \top -skew-Hamiltonian matrices $\widehat{\mathcal{K}}$, $\widehat{\mathcal{N}}$, starting vector y_1 and shift value τ .

Output: \mathfrak{m} eigenpairs of $(\widehat{\mathcal{K}}, \widehat{\mathcal{N}})$.

1: Use Algorithm 5 with starting vector y_1 to generate a generalized \top -isotropic Arnoldi decomposition with order \mathfrak{m} : $\widehat{\mathcal{K}}Z_{\mathfrak{m}} = Y_{\mathfrak{m}}H_{\mathfrak{m}} + h_{\mathfrak{m}+1,\mathfrak{m}}y_{\mathfrak{m}+1}e_{\mathfrak{m}}^{\top},$

 $\widehat{\mathcal{N}}Z_{\mathfrak{m}} = Y_{\mathfrak{m}}R_{\mathfrak{m}}.$

2: repeat

- 3: Use Algorithm 5 to extend the generalized \top -isotropic Arnoldi decomposition with order m to order $(\mathfrak{m} + p)$: $\widehat{\mathcal{K}Z}_{\mathfrak{m}+p} = Y_{\mathfrak{m}+p}H_{\mathfrak{m}+p} + h_{\mathfrak{m}+p+1,\mathfrak{m}+p}y_{\mathfrak{m}+p+1}e_{\mathfrak{m}+p}^{\top},$
 - $\widehat{\mathcal{N}}Z_{\mathfrak{m}+p} = Y_{\mathfrak{m}+p}R_{\mathfrak{m}+p}.$
- 4: Use Krylov–Schur restarting scheme [20,21] to reform a new generalized ⊤-isotropic Arnoldi decomposition with order m.
- 5: **until** wanted \mathfrak{m} eigenpairs of $(\widehat{\mathcal{K}}, \widehat{\mathcal{N}})$ are convergent

steps just mentioned are Step 5 and Step 14, respectively, in Algorithm 5. Moreover, the GTSHIRA algorithm based on the generalized \top -isotropic Arnoldi process is presented in Algorithm 6 for finding eigenpairs of the matrix pair $(\widehat{\mathcal{K}}, \widehat{\mathcal{N}})$.

In the above TSHIRA and GTSHIRA algorithms, the main costs arise in computing $u_{j+1} = \mathcal{B}u_j$ and solving linear system $\widehat{\mathcal{N}}z_j = y_j$ at the *j*th \top -isotropic and generalized \top -isotropic Arnoldi steps, respectively. From (14b), (15) and (16), computing these vectors u_{j+1} and z_j require to solve the following linear systems

$$\mathcal{N}_1 v_1 = b_1, \qquad \mathcal{N}_2 v_2 = b_2. \tag{23}$$

By the definitions of \mathcal{M} and \mathcal{L} in (11), we see that solving (23) is equivalent to solve

$$(\tau^2 A_1^{\top} + \tau A_0 + A_1) v_{11} = b_{11} - \tau b_{12}, \tag{24}$$

$$v_{12} = -b_{12} - (A_0 + \tau A_1^{\top})v_{11},$$

and

$$(\tau^2 A_1 + \tau A_0 + A_1^{\top}) v_{22} = b_{22} + (A_0 + \tau A_1) b_{21},$$
(25)

 $v_{21} = \tau v_{22} - b_{21},$

where $v_1 = [v_{11}^{\top}, v_{12}^{\top}]^{\top}$, $v_2 = [v_{21}^{\top}, v_{22}^{\top}]^{\top}$, $b_1 = [b_{11}^{\top}, b_{12}^{\top}]^{\top}$ and $b_2 = [b_{21}^{\top}, b_{22}^{\top}]^{\top}$. By the definitions of A_0 and A_1 , it holds that $\tau^2 A_1^{\top} + \tau A_0 + A_1 = (G + \tau F)M_2^{-1}(F^{\top} + \tau G^{\top}) - \tau M_1$ (26)

Algorithm 7 GE_GTSHIRA/GE_TSHIRA

Input: matrices *F*, *G*, M_2 and M_1 , shift value τ and the number m of desired eigenvalues.

Output: eigenpairs $\{(\gamma_j, [(\psi_{i,j}^{(1)})^{\top}, (\psi_{\ell,j}^{(1)})^{\top}]^{\top}), (\gamma_j^{-1}, [(\psi_{i,j}^{(2)})^{\top}, (\psi_{\ell,j}^{(2)})^{\top}]^{\top})\}_{j=1}^{\mathfrak{m}}$ of (1) where $\gamma_j + \gamma_j^{-1}$ for $j = 1, ..., \mathfrak{m}$ are the closest to shift value $\tau + \tau^{-1}$.

- 1: Compute eigenpairs $\{(\widehat{\mu}_j, z_j \equiv [z_{j1}^{\top}, z_{j2}^{\top}]^{\top})\}_{j=1}^{\mathfrak{m}}$ of $(\widehat{\mathcal{K}}, \widehat{\mathcal{N}})$ by using GTSHIRA or Compute eigenpairs $\{(\widehat{\mu}_j, \widetilde{z}_j)\}_{j=1}^{\mathfrak{m}}$ of \mathcal{B} by using TSHIRA and solve $\mathcal{N}_2[z_{j1}^{\top}, z_{j2}^{\top}]^{\top} = \widetilde{z}_j$, for $j = 1, \ldots, \mathfrak{m}$.
- 2: Compute eigenvalues γ_i and γ_i^{-1} of TPQEP in (2) by solving

$$\gamma^2 - (\tau + \tau^{-1} + \widehat{\mu}_j^{-1})\gamma + 1 = 0;$$

Compute eigenvectors

$$\psi_{i,j}^{(1)} \equiv \gamma_j^{-1} z_{j1} - z_{j2}, \quad \psi_{i,j}^{(2)} \equiv \gamma_j z_{j1} - z_{j2}$$

corresponding to γ_j , γ_j^{-1} , respectively, for $j = 1, 2, ..., \mathfrak{m}$.

3: Compute

$$\psi_{\ell,j}^{(1)} = -M_2^{-1} \left(\gamma_j^{-1} F^\top \psi_{i,j}^{(1)} + G^\top \psi_{i,j}^{(1)} \right), \ \psi_{\ell,j}^{(2)} = -M_2^{-1} \left(\gamma_j F^\top \psi_{i,j}^{(2)} + G^\top \psi_{i,j}^{(2)} \right)$$

for j = 1, ..., m.

and

$$\tau^2 A_1 + \tau A_0 + A_1^{\top} = (F + \tau G) M_2^{-1} (G^{\top} + \tau F^{\top}) - \tau M_1.$$
(27)

Let $M_1 = LU$ be the LU factorization of M_1 and set

$$E_1 = L^{-1}\left(\frac{1}{\tau}G + F\right), \qquad E_2 = U^{-\top}(F + \tau G).$$
 (28)

By the Sherman-Morrison-Woodbury formula, (26) and (27) imply that

$$\left(\tau^{2}A_{1}^{\top}+\tau A_{0}+A_{1}\right)^{-1}=-\frac{1}{\tau}U^{-1}\left[I+E_{1}\left(M_{2}-E_{2}^{\top}E_{1}\right)^{-1}E_{2}^{\top}\right]L^{-1}$$

and

$$\left(\tau^{2}A_{1}+\tau A_{0}+A_{1}^{\top}\right)^{-1}=-\frac{1}{\tau}L^{-\top}\left[I+E_{2}\left(M_{2}-E_{1}^{\top}E_{2}\right)^{-1}E_{1}^{\top}\right]U^{-\top},$$

respectively.

Obviously, from (28), we need *m* forward substitutions and *m* backward substitutions to obtain E_1 and E_2 , respectively. Furthermore, in addition to the cost in solving small linear systems $(M_2 - E_2^{\top}E_1)^{-1}$ and $(M_2 - E_1^{\top}E_2)^{-1}$, only two forward substitutions $(L^{-1}, U^{-\top})$ and two backward substitutions $(L^{-\top}, U^{-1})$ are required to obtain the solutions of (24) and (25) for generating Krylov subspace at each iterative step. Recall that, for GE_SDA and GE_SA, in order to form the matrices A_0 and A_1 in (3), one needs to compute $M_1^{-1}F$ and $M_1^{-1}G$ which require 2*m* forward and backward substitutions. Since the shift-and-invert Arnoldi method is known to converge very fast when a proper shift is known, the overall computational costs of GE_GTSHIRA and GE_TSHIRA, including computing E_1, E_2 and solving linear systems in each iterative steps, can be only about half amount of the computation cost needed in GE_SDA and GE_SA. Our numerical results in Table 2 confirm this observation. Finally, we summarize the process of applying TSHIRA/GTSHIRA to solve the GEP in (1) in Algorithm 7 and show the comparison of the computational costs for TSHIRA and GTSHIRA in Table 2.

4. Numerical results

In this section, we tests the above mentioned four types of structure preserving algorithms on computing the dispersion diagram of the frequency that are close to the stopping frequency of the SAW filter. The piezoelectric substrate of the filter is made of 15° rotated quartz. The configuration of our computational domain is shown in Fig. 1 where the domain width \overline{AB} and height \overline{CD} are set to be 10^{-6} and 3×10^{-6} , respectively, the ratio of the electrode width \overline{EF} versus the domain width is set to be $\frac{1}{2}$ and the ratio of the electrode thickness \overline{DE} versus the domain height is $\frac{1}{15}$. In our numerical studies, the viscous damping coefficient κ_1 is set to be 10^{-14} and the mass damping coefficient κ_2 is taken as $1 - \kappa_1$ to account for the effect from the electrode weight. All computations are carried out in MATLAB 2010b on a HP workstation with an Intel Quad-Core Xeon X5570 2.93 GHz and 60 GB main memory, using IEEE double-precision floating-point arithmetic.

Suppose m reciprocal pairs of eigenvalues near \mathbb{U} are desired. For TSHIRA and GTSHIRA, the restart procedure will be activated when the desired eigenpairs do not converge before the dimension of the Krylov subspace reaches 5m. This is done by setting the value of *p* in Step 3 of Algorithms 4 and 6 to 4m. In the following discussion, we take m = 5 and the

| | | TSHIRA | GTSHIRA |
|--------------------|---|------------------------|------------------------|
| Compute E_1, E_2 | $M_1 = LU$ | 1 | 1 |
| | $F + \xi G$ | 2 | 2 |
| | Solve $Lx = b_1$ | т | т |
| | Solve $U^{\top}y = c_2$ | т | т |
| | $E_2^{\top}E_1$ (flops) | $2m^2n$ | $2m^2n$ |
| jth step Arnoldi | Solve $Lx = b_1, L^{\top}y = c_1$ | 1 | 1 |
| | Solve $Ux = b_2$, $U^{\top}y = c_2$ | 1 | 1 |
| | Compute Fd_1 , $F^{\top}c_1$, Gd_2 , $G^{\top}c_2$ | 3 | 3 |
| | Compute M ₁ b | 2 | 2 |
| | Compute $E_1d_1, E_1^{\top}c_1, E_2d_2, E_2^{\top}c_2$ | 1 | 1 |
| | Saxpy and inner products (flops) | 8nj + 15n | 16nj + 18n |
| Schur restarting | Matrix product (flops) | $2(\mathfrak{m}+p)^2n$ | $4(\mathfrak{m}+p)^2n$ |

 Table 2

 Computational costs for TSHIRA and GTSHIRA.



Fig. 2. The distribution of the eigenvalues which are close to and inside of \mathbb{U} .



Fig. 3. Dispersion diagrams of α and β near the stopping band.

matrix dimensions of C_i and C_b are $n = 63\,960$ and m = 723, respectively. An example of computed reciprocal eigenpairs near U at frequency $\omega = 1.2757/(2\pi) \times 10^{10}$ is shown in Fig. 2. The dispersion diagrams of the attenuation constant α and the propagation constant β associated with the eigenvalue $\lambda(\omega)$ are shown in Fig. 3, for frequency ω around the stopping band, where the eigenpair most close to -1 on the complex plane is plotted.

4.1. Accuracy of structure-preserving eigensolvers

In this subsection, we compare the accuracy of the eigenpairs, computed by structure-preserving Algorithms 1, 2 and 7, respectively, for the GEP (8). Recall that the Krylov subspace \mathcal{U}_j generated by the \top -Hamiltonian matrix \mathcal{B} is automatically \top -isotropic in Theorem 3.2, and the subspaces \mathcal{Z}_j and \mathcal{Y}_{j+1} generated in Theorem 3.4 are automatically \top -bi-isotropic. As mentioned in Sections 3.3 and 3.4, isotropic re-orthogonalization in Step 6 of Algorithm 3 and Steps 7 and 16 of Algorithm

Table 3

Convergent eigenvalues computed by T_NoSymp and GT_NoBilso at $\omega = 1.2757/(2\pi) \times 10^{10}$.

| | T_NoSymp | GT_NoBiIso |
|---|---|---|
| | -0.85175542558 - 0.52335156640i -0.85228028786 + 0.52367406214i | $\begin{array}{c} -0.85175542559 - 0.52335156640\imath \\ -0.85228028785 + 0.52367406213\imath \end{array}$ |
| $\left(\lambda, \frac{1}{\lambda}\right)$ | -0.85175542557 - 0.52335156639ı -0.85228028787 + 0.52367406214ı | $\begin{array}{l} -0.85175542556-0.52335156641 \imath \\ -0.85228028786+0.52367406216 \imath \end{array}$ |
| (, , / | $\begin{array}{l} -0.98999503056 + 0.00448884999 \imath \\ -1.01008531402 - 0.00457994365 \imath \end{array}$ | $\begin{array}{l} -0.98999503056+0.00448884999\imath\\ -1.01008531402-0.00457994365\imath\end{array}$ |
| | -0.98999503056 + 0.00448884999i -1.01008531402 - 0.00457994365i | -0.98999503056 + 0.00448884999i -1.01008531402 - 0.00457994365i |



Fig. 4. The relative residual of the computed eigenpairs produced by different re-bi-isotropic processes in Algorithm 5 with shift value $\tau = -0.99$.

5 is important in maintaining the \top -isotropic property. Moreover, Theorems 3.3 and 3.5 both show that the multiplicities of eigenvalues of $(\mathcal{K}, \mathcal{N})$ are all even. In other words, no duplicate eigenpairs need to be computed theoretically when the \top -isotropic property is kept during the computation. On the other hand, without the isotropic re-orthogonalization process, extra computation cost can arise in computing the duplicate eigenpairs. We would like to address this issue by numerical studies shown in the following. We also like to point out that the accuracy of the computed eigenpairs can be affected by different approaches in isotropic re-orthogonalization.

First, let us denote the algorithm that applying TSHIRA without the re-symplectic process as T_NoSymp and the algorithm that applying GTSHIRA without these re-bi-isotropic processes as GT_NoBiIso. In Table 3, the convergent eigenvalues obtained by T_NoSymp and GT_NoBiIso at frequency $\omega = 1.2757/(2\pi) \times 10^{10}$ are listed. Obviously, one can see that, in case only two eigenpairs { $(\lambda_1, \lambda_1^{-1}), (\lambda_2, \lambda_2^{-1})$ } are needed here, the algorithms T_NoSymp and GT_NoBiIso return four convergent eigenpairs in which two of them are indeed the duplicated pairs.

Next, let us compare the accuracy of the computed eigenpairs obtained from three different isotropic reorthogonalization approaches in GTSHIRA. One or two steps of re-bi-isotropic process can be performed by the for-loops in Steps 6–8 and 15–17.

To distinguish among various re-bi-isotropic processes, we use notations "FullIso", "zIsoY" and "yIsoZ" defined as follows:

- FullIso: Algorithm 5 with two for-loops in Steps 6–8 and 15–17.
- zIsoY: Algorithm 5 with one for-loop in Steps 6–8 and omitting for-loop in Steps 15–17.
- yIsoZ: Algorithm 5 with one for-loop in Steps 15–17 and omitting for-loop in Steps 6–8.

To measure the accuracy of computed eigenpairs of (8), we consider the relative residual of an eigenpair (λ, ψ) where $\psi = [\psi_i^{\top}, \psi_{\rho}^{\top}]^{\top}$ which is defined as following:

$$\frac{\left\| \begin{bmatrix} C_{i} & C_{i\ell} \\ C_{ir}^{\top} & 0 \end{bmatrix} \psi - \lambda \begin{bmatrix} 0 & C_{ir} \\ C_{i\ell}^{\top} & C_{b} \end{bmatrix} \psi \right\|_{F}}{\left\| \begin{bmatrix} C_{i} & C_{i\ell} \\ C_{ir}^{\top} & 0 \end{bmatrix} \right\|_{F} \|\psi\|_{F} + |\lambda| \left\| \begin{bmatrix} 0 & C_{ir} \\ C_{i\ell}^{\top} & C_{b} \end{bmatrix} \right\|_{F} \|\psi\|_{F}},$$

here $||*||_F$ is the Frobenius matrix norm. The relative residuals of the convergent eigenpairs computed by "FullIso", "zIsoY" and "yIsoZ" are shown in Fig. 4. From the numerical results in Fig. 4, we see that the accuracy of the convergent eigenpairs



Fig. 5. The errors of the equalities in (18) and (19) for "FullIso", "zIsoY" and "yIsoZ".

| | TSHIRA | GTSHIRA | SDA | SA |
|---|--------|---------|--------|--------|
| Compute $C_i = LU$ | 191.31 | 191.31 | 191.31 | 191.31 |
| Compute $E_1, E_2, E_2^{\top} E_1$ | 243.75 | 243.75 | | |
| Compute A_0, A_1 | | | 533.94 | 533.94 |
| Solve dense TPQEP | | | 34.145 | |
| Solve $Lx = b_1$ | 4.9850 | 4.9850 | 1.9940 | 1.9940 |
| Solve $Ux = b_2$ | 3.9775 | 3.9775 | 1.5910 | 1.5910 |
| Solve $U^{\top}y = c_2$ | 33.597 | 27.998 | | |
| Solve $L^{\top}y = c_1$ | 36.300 | 30.250 | | |
| Compute $E_1d_1, E_2^{\top}c_2$ | 4.9150 | 4.9150 | | |
| Compute $E_1^{\top}c_1, \tilde{E_2}d_2$ | 5.7930 | 4.8275 | | |

computed by "yIsoZ" is higher than those by "FullIso" and "zIsoY". This result can be explained from the accumulation of the errors in the equalities (18) and (19). Let $\xi_{j,K} \equiv \|\widehat{\mathcal{K}}Z_j - Y_j\widehat{H}_j - \widehat{h}_{j+1,j}y_{j+1}e_j^\top\|_2$ and $\xi_{j,N} \equiv \|\widehat{\mathcal{K}}Z_j - Y_j\widehat{R}_j\|_2$, denote these errors in the *j*th iteration. The error $\xi_{j,N}$ depends on the accuracy of the solution of the linear systems in (23). If z_j is reorthogonalized to $\mathscr{J}\overline{Y}_j$, then the error produced by this reorthogonalization will reduce the accuracy of $\xi_{j,N}$. Therefore, $\xi_{j,N}$ produced by "FullIso" and "zIsoY" are greater than that by "yIsoZ" as shown in Fig. 5(a). On the other hand, the error $\xi_{j,K}$ only depends on the accuracy of matrix product vector and vector inner product. Obviously, the amount of $\xi_{j,K}$ is much less than the amount of $\xi_{j,N}$. Consequently, even though the accuracy of $\xi_{j,K}$ can reduced by the errors from reorthogonalization y_{j+1} to $\mathscr{J}\overline{Z}_j$ as shown Fig. 5(b), the reorthogonalization process "yIsoZ" is much accurate than the "FullIso" and "zIsoY" reorthogonalization process "yIsoZ" is much accurate than the "FullIso" and "zIsoY" reorthogonalization process "yIsoZ" is much accurate than the "FullIso" and "zIsoY" reorthogonalization processes.

Finally, we compare the accuracy of the eigenpairs $(\lambda(\omega), u(\omega))$ obtained from GE_SDA, GE_SA, GE_TSHIRA and GE_GTSHIRA with "ylsoZ" re-bi-isotropic process. The relative residuals resulted from these algorithms in computing four reciprocal eigenpairs $(\lambda_i(\omega), u_i(\omega))$, for i = 1, ..., 4, that are closest to -1 on the complex plane are plotted in Fig. 6 for each frequency ω near the stopping band. Obviously, one can see that the accuracy of the eigenpairs obtained from GE_SDA and GE_SA are higher than those obtained by GE_TSHIRA and GE_GTSHIRA.

4.2. Comparison with computational costs

In this subsection, we discuss the computational costs of structure-preserving Algorithms 1, 2 and 7 in computing m = 5 desired eigenpairs. Our numerical results show that the desired eigenpairs are convergent within 5m-isotropic Arnoldi steps without restart for GE_TSHIRA and GE_GTSHIRA. On the other hand, it requires total 18 iterations to obtain a convergent X_k in Steps 3–6 for the SDA algorithm. As we mentioned in Section 3.3, the number of forward and backward substitutions needed for GE_TSHIRA and GE_GTSHIRA is only about half the amount of these substitutions that needed to transform the GEP into TPQEP in GE_SDA and GE_SA. Since only additional 25 forward substitutions and backward substitutions are needed in GE_TSHIRA and GE_GTSHIRA for solving linear systems Lx = b and Uy = c, we expect GE_TSHIRA and GE_SDA. The following numerical results support this observation.

To give an overall comparison for GE_SDA, GE_SP, GE_TSHIRA and GE_GTSHIRA, in Table 4, computational intensive items in these algorithms are listed in the first column and the sums of the CPU times for each associated item are listed in the other four columns. From the results in Table 4, the dominant computational costs in GE_TSHIRA and GE_GTSHIRA are the costs for computing E_1 , E_2 , $E_2^{\top}E_1$ and LU factorization of C_i . For GE_SDA and GE_SA, the cost in computing the matrices A_0 and A_1 of the TPQEP is the main cost comparing to the other costs. Obviously, the numbers shown in Table 4 indicate that GE_TSHIRA and GE_GTSHIRA are more efficient than GE_SDA and GE_SA. We also plot the overall CPU times for GE_SDA



Fig. 6. Relative residuals for GE_SDA, GE_SA, GE_TSHIRA and GE_GTSHIRA with shift value $\tau = -0.89$.



Fig. 7. CPU times for GE_SDA and GE_GTSHIRA.

and GE_GTSHIRA with frequency from $1.274/(2\pi) \times 10^{10}$ to $1.279/(2\pi) \times 10^{10}$ in Fig. 7. From Fig. 7, one can see that the total CPU times needed in GE_SDA and GE_SA are 40% more than the CPU time needed in GE_TSHIRA and GE_GTSHIRA for computing 5 desired eigenpairs.

5. Conclusion

In this paper, we have discussed the structure-preserving methods for solving the generalized eigenvalue problem arising in the surface acoustic wave propagation on a simple resonator with an interdigital transducer (IDT) where electrodes are arranged periodically on piezoelectric substrates (PZT) such as 15° rotated Quartz. With given periodic boundary conditions,

the eigenvalues of the GEP appear in the reciprocal pairs (λ , λ^{-1}). In order to preserve the reciprocal relationship of the eigenvalues, the GEP is transformed to two types of T-palindromic quadratic eigenvalue problems, one with large coefficient matrices and the other with small coefficient matrices. The structure-preserving algorithms GE_SDA and GE_SA in Algorithms 1 and 2 are employed to solve the TPQEP (3) with small-size coefficient matrices and GE_TSHIRA and GE_GTSHIRA in Algorithm 7 are employed to solve the TPQEP (4) with large-size coefficient matrices.

In finding the five eigenpairs that are near \mathbb{U} and close to -1, we observed duplicate eigenpairs appear when applying GE_TSHIRA and GE_GTSHIRA without re-symplectic and re-bi-isotropic processes, respectively. On the other hand, no duplicate eigenpairs are observed when re-symplectic and re-bi-isotropic processes are integrated in GE_TSHIRA and GE_GTSHIRA. Three different re-bi-isotropic processes in GE_GTSHIRA has been tested. We have found that using the re-bi-isotropic processes in Steps 15–17 of Algorithm 5 achieves the best accuracy. Moreover, our numerical results show that the relative residuals of the eigenpairs produced by GE_SDA/GE_SA and GE_TSHIRA/GE_GTSHIRA can be less than 10^{-17} and 10^{-15} , respectively. Although the accuracy of GE_SDA and GE_SA is marginally higher than that of GE_TSHIRA and GE_GTSHIRA, we further found that the total CPU times required for computing the five desired eigenpairs by GE_SDA and GE_SA are about 40% more than that are required by GE_TSHIRA and GE_GTSHIRA. Therefore, by transforming the GEP into the TPQEP (4), the structure-preserving Arnoldi type algorithm GE_TSHIRA or GE_GTSHIRA with one "re-symplectic" or "re-bi-isotropic" processes provide an accurate and efficient way in finding the reciprocal eigenpairs of the GEP (1).

For further reading

[21,22].

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