Computation of scattering resonances in absorptive and dispersive media with applications to metal-dielectric nano-structures

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Abstract
In this paper we consider scattering resonance computations in optics when the resonators consist of frequency dependent and lossy materials, such as metals at optical frequencies. The proposed computational approach combines a novel $hp$-FEM strategy, based on dispersion analysis for complex frequencies, with a fast implementation of the nonlinear eigenvalue solver NLEIGS. Numerical computations illustrate that the pre-asymptotic phase is significantly reduced compared to standard uniform $h$ and $p$ strategies. Moreover, the efficiency grows with the refractive index contrast, which makes the new strategy highly attractive for metal-dielectric structures. The $hp$-refinement strategy together with the efficient parallel code result in highly accurate approximations and short runtimes on multi processor platforms.

Keywords:
Plasmon resonance, Resonance modes, Nonlinear eigenvalue problems, Helmholtz problem, PML, Dispersion analysis, leaky modes, resonant states, quasimodes, quasi-normal modes

1. Introduction

Metallic nano-structures play an important role in many applications in physics, including surface enhanced Raman scattering and optical antennas [1]. Surface plasmons that may exist in these structures cause an enormous electromagnetic field enhancement near the surface of noble metals. In nanomedicine gold nanoparticles are used in the forefront of cancer research since they not only support plasmon resonances but also have excellent biocompatibility [2].

The material properties of metals are characterized by the complex relative permittivity function $\varepsilon$, which changes rapidly at optical frequencies $\omega$. The most common accurate material model is then the Drude-Lorentz model

$$\varepsilon_{\text{metal}}(\omega) := \varepsilon_\infty + \sum_{j=0}^{N_p} f_j \omega_p^2 \left( \frac{\omega_j^2}{\omega_j^2 - \omega^2 - i\omega \gamma_j} \right),$$

where $\varepsilon_\infty \geq 1$ and $f_j$, $\omega_p$, $\omega_j$, $\gamma_j$ are non-negative [3]. Hence, the Maxwell eigenvalue problem in the spectral parameter $\omega$ is nonlinear for metal-dielectric nanostructures. Research in operator...
theory for this type of non-selfadjoint operator functions is in its infancy and has been focused on photonic crystal applications [4].

In this article, we consider open systems in nano-optics, where the material properties are modeled by (1). The most common approach to characterize the optical properties of open metal-dielectric nanostructures is to solve a source problem in time-domain and search for peaks in the amplitude of the field [5]. Another common strategy is to solve a source problem for a fixed real frequency and perform a frequency sweep in a region of interest [6]. These two strategies give valuable information of the structure for a given source. A highly attractive alternative that is used in this paper is to characterize the behavior of the system using scattering resonances [7, 8].

Resonances are solutions to a nonlinear eigenvalue problem with a Dirichlet-to-Neumann map (DtN) on an artificial boundary [9, 10]. An attractive alternative is to use a perfectly matched layer (PML) [11, 12]. This method was introduced for source problems in electromagnetics by Berenger [13] and it is related to complex coordinate stretching developed in quantum mechanics [14, Chapter 16]. The application of the PML method for resonance problems has the advantage that for non-dispersive refractive indices the resulting matrix eigenvalue problem is linear, and the eigenvalue problem is rational when a Drude-Lorentz model is used. Resonance computations with a dispersive refractive index are demanding since nonphysical eigenvalues may appear in the region of interest if the approximation properties of the used finite element space are not very good; See [15] for a discussion of spurious eigenvalues in the one dimensional case.

The linear algebra problem that must be solved in this kind of computations is a rational eigenvalue problem, a particular case of the nonlinear eigenvalue problem \( T(\omega)\xi = 0 \). Recently, several numerical methods have been proposed to compute a few eigenvalues \( \omega \) (and corresponding eigenvectors \( \xi \)) of large-scale nonlinear eigenvalue problems [16, 17, 18]. Some of these methods are available in the SLEPc library [19]. Essentially, there are three types of methods: Newton-type methods, contour integral methods, and linearization methods. Newton-type methods rely on having a good initial guess, otherwise the iteration may converge to an eigenvalue far from the search region. Contour integral methods compute all eigenvalues contained in a prescribed region of the complex plane, but they require having a good estimate of the number of enclosed eigenvalues, and on the other hand they have a high computational cost since they require a matrix factorization at each integration point. In this paper, we consider a method of linearization type, namely NLEIGS, see §4.5.

2. Resonances in optical nano-structures

Our aim is to compute resonances in nano-structures using accurate material models for e.g. metals at optical frequencies. This requires a permittivity function \( \varepsilon \) that depends on the spectral parameter \( \omega \). For many metals, the real part of \( \varepsilon \) is negative in the optical region, which is explored
in plasmonics [1]. Below, we state well known properties for isotropic passive materials that are valid for any fixed $x \in \mathbb{R}^d$. Let

$$C_+ := \{ z \in \mathbb{C} : 0 \leq \arg z < \pi, z \neq 0 \}.$$ 

Then $\omega \varepsilon(\omega) \in C_+$ for $\omega \in C_+$, where $\varepsilon$ never vanishes in $\mathbb{C}_+$ [3]. The most common material model for solid materials such as Gold, Silver, and Silica is the Drude-Lorentz (1) model. This rational model of $\omega$ satisfies the stated analytical requirements and will be used in the applications part of the article.

Assume that $\varepsilon(x, \omega) = \varepsilon(x_1, x_2, \omega)$ is independent of $x_3$ and consider waves propagating in the $(x_1, x_2)$-plane. The $x_3$-independent electromagnetic field $(E, H)$ is then decomposed into transverse electric (TE) polarized waves $(E_1, E_2, 0, 0, H_3)$ and transverse magnetic (TM) polarized waves $(0, 0, E_3, H_1, H_2, 0)$ [3]. This decomposition reduces Maxwell’s equations to one scalar equation for $H_3$ and one scalar equation for $E_3$. The TM-polarized waves and the TE-polarized waves satisfies formally

$$-\Delta E_3 - \omega^2 \varepsilon E_3 = 0 \quad \text{and} \quad -\nabla \cdot \left( \frac{1}{\varepsilon} \nabla H_3 \right) - \omega^2 H_3 = 0,$$

respectively. The full vector fields $(E, H)$ are then obtained from Maxwell’s equations

$$E = \frac{-1}{i \omega \varepsilon(\omega)} \nabla \times H, \quad \text{and} \quad H = \frac{1}{i \omega} \nabla \times E.$$  \(3\)

For simplicity, we consider first resonances in the TM-case with an $\omega$-independent permittivity function $\varepsilon \geq 1$, where $\varepsilon - 1$ has compact support. Let $L^2_{\text{comp}}$ denote the space of $L^2$-functions vanishing outside some compact set and let $L^2_{\text{loc}}$ denote the space of functions that are in $L^2(K)$ for every compact subset $K$ of $\mathbb{R}^d$. Define the operator $A : L^2(\mathbb{R}^d) \to L^2(\mathbb{R}^d)$ with domain $\text{dom} A = H^2(\mathbb{R}^d)$ by $Au := -\varepsilon^{-1} \Delta u$. The spectrum $[0, \infty)$ is then continuous [14] and we denote by $R(\omega) : L^2(\mathbb{R}^d) \to L^2(\mathbb{R}^d)$ the resolvent

$$R(\omega) := (A - \omega^2)^{-1}, \quad \text{Im} \omega^2 > 0.$$ 

The operator function $R$ is a meromorphic family of operators that can be extended to

$$\hat{R}(\omega) : L^2_{\text{comp}}(\mathbb{R}^d) \to L^2_{\text{loc}}(\mathbb{R}^d), \quad \hat{R}(\omega) := (A - \omega^2)^{-1}, \quad \text{Im} \omega^2 > 0.$$ 

The scattering resonances are then defined as the poles of the meromorphic continuation of $\hat{R}$ to $\mathbb{C}$. The functions in $L^2_{\text{loc}}(\mathbb{R}^d)$ that correspond to a scattering resonance are called resonance modes [7]. Note that for metal-dielectric nanostructures the operator $A$ in the TM-case is replaced with an operator function in $\omega$. In the next sections, we will describe two common approaches to compute resonances and the restriction of resonance modes to a compact subset of $\mathbb{R}^d$. In the following, we use the notation

$$-\nabla \cdot \left( \rho \nabla u \right) - \omega^2 \eta u = 0,$$

where $u := E_z$, $\rho := 1$, $\eta := \varepsilon$ for the TM-case and $u := H_z$, $\rho := 1/\varepsilon$, $\eta := 1$ for the TE-case.
2.1. Scattering resonances in $\mathbb{R}$

Resonances as discussed in section 2 can also be determined from a problem with a Dirichlet-to-Neumann (DtN) map [9, 20]. In one space dimension the resonance problem restricted to $I_a := (-a, a)$ is formerly: Find a non-zero $u$ and a complex $\omega$ such that

$$-(\rho u')' - \omega^2 \eta u = 0 \text{ for } x \in I_a,$$

where the Dirichlet-to-Neumann (DtN) map at $x = \pm a$ is

$$u'(-a) = -i\omega u(-a), \quad u'(a) = i\omega u(a).$$

Let $\mathcal{Z}$ denote the set of values $\omega$ that are zeros or poles of $\varepsilon$ and set $\mathcal{D} := \mathbb{C} \setminus \mathcal{Z}$. Define for $u, v \in H^1(I_a)$ and $\omega \in \mathcal{D} \subset \mathbb{C}$ the forms

$$\tilde{t}_0(\omega)[u, v] := \int_{-a}^{a} \rho uu' dx, \quad \tilde{t}_1(\omega)[u, v] := -iu(a)v(a) - iu(-a)v(-a), \quad \tilde{t}_2(\omega)[u, v] := -\int_{-a}^{a} \eta uv dx.$$

The nonlinear eigenvalue problem is then as follows: Find vectors $u \in H^1(I_a) \setminus \{0\}$ and $\omega \in \mathcal{D}$ satisfying

$$\tilde{t}_1(\omega)[u, v] := \omega^2 \tilde{t}_2(\omega)[u, v] + \omega \tilde{t}_1(\omega)[u, v] + \tilde{t}_0(\omega)[u, v] = 0$$

for all $v \in H^1(I_a)$.

Let $I_a = I_0 \cup \ldots \cup I_N$, denote a partitioning of $I_a$ and $\chi_{I_m}$ the characteristic function of the subset $I_m$. For material properties that are piecewise constant in $x$, we assume a permittivity function in the form

$$\varepsilon(x, \omega) := \sum_{m=0}^{N} \varepsilon_m(\omega) \chi_{I_m}(x), \quad x \in I_a, \quad \omega \in \mathcal{D},$$

where the dependencies on $\omega$ in $\varepsilon_m$ for $m = 0, 1, \ldots$ are of Drude-Lorentz type (1). Note that (7) is an quadratic eigenvalue problem if $\varepsilon$ is independent of $\omega$ and a rational eigenvalue problem for Drude-Lorentz type of materials.

2.2. Scattering resonances in $\mathbb{R}^2$

Resonances in $\mathbb{R}^2$ can also be approximated by (4) with a DtN-map on an artificial boundary [9, 10]. However, the nonlinearity in the DtN-map is more complicated in dimensions larger than one. Then, an attractive alternative to the DtN-map is a complex coordinate stretching technique called Perfectly Matched Layers (PML). This approach does not add any non-linearity to the problem. Hence, in our setting we will obtain a rational eigenvalue problem. Approximation of resonances using a radial PML was analyzed in [11] and we consider the truncation of the infinite PML problem to the disc $\Omega$ in $\mathbb{R}^2$.

Let $\Omega_a$ denote a disk of radius $a$, and let $\Omega_1, \Omega_2, \ldots, \Omega_N$, denote the subsets of $\Omega_a$ corresponding to the resonators. Set $\Omega_r := \cup_{i=1}^{N} \Omega_i$, $\Omega_0 := \Omega_a \setminus \Omega_r$, and attached to $\Omega_a$ an outer layer $\Omega_{PML}$. Then, the computational domain is the disc $\Omega := \Omega_a \cup \Omega_{PML}$ as illustrated in Fig. 1.

We define the complex stretching functions in polar coordinates $(r, \theta)$, similarly as presented in [21, 11]:

$$\tilde{\sigma}(r) := \begin{cases} 0, & \text{if } r < a \\ P(r), & \text{if } a \leq r \leq b, \\ \sigma_0, & \text{if } r > b \end{cases}, \quad \tilde{\alpha}(r) := 1 + i\tilde{\sigma}(r), \quad \tilde{\tau}(r) := (1 + i\tilde{\sigma})r = \tilde{\alpha}(r)r,$$

$$\sigma(r) := \tilde{\sigma}(r) + r \frac{\partial \tilde{\sigma}}{\partial r}, \quad \alpha(r) := \frac{\partial \tilde{\tau}}{\partial r} = 1 + i\sigma(r).$$
where the polynomial $P(r)$ is required to be increasing in $[a, b]$, and $\tilde{\sigma}(r) \in C^2(0, \ell)$, $\sigma(r) = \partial(r\tilde{\sigma})/\partial r$. For this we introduce the fifth order polynomial $P(r)$ satisfying $P(a) = P'(a) = P''(a) = P'(b) = P''(b) = 0$ and $P(b) = \sigma_0$.

From the given curved coordinate representation, we transform to Cartesian coordinates and define $A \in C^2(\Omega)^{2 \times 2}$ and $B \in C^2(\Omega)$ by

$$A := \begin{pmatrix} \frac{a}{\alpha} \cos^2 \theta + \frac{a}{\alpha} \sin^2 \theta & \left( \frac{a}{\alpha} - \frac{a}{\alpha} \right) \sin \theta \cos \theta \\ \left( \frac{a}{\alpha} - \frac{a}{\alpha} \right) \sin \theta \cos \theta & \frac{a}{\alpha} \sin^2 \theta + \frac{a}{\alpha} \cos^2 \theta \end{pmatrix}, \quad B := \alpha \tilde{\alpha}.$$  

(10)

The PML coefficients are illustrated in Fig. 1, and it can be seen that $A$ and $B$ are identities for $r \leq a$.

Let $(\cdot, \cdot)_{\Omega_j}$ denote the inner product in $L^2(\Omega_j)$. The nonlinear eigenvalue problem is then: Find $u \in H^1_0(\Omega) \setminus \{0\}$ and $\omega \in \mathcal{D}$ such that for all $v \in H^1_0(\Omega)$

$$t_2(\omega)[u, v] = 0,$$

(11)

where $t_2(\omega)[u, v] := \tilde{t}_0(\omega)[u, v] + \tilde{t}_1(\omega)[u, v]$ with

$$\tilde{t}_0(\omega)[u, v] := (\rho \nabla u, \nabla v)_{\Omega_a} - \omega^2 (\rho u, v)_{\Omega_a}, \quad \tilde{t}_1(\omega)[u, v] := (A \nabla u, \nabla v)_{\Omega_{PML}} - \omega^2 (B u, v)_{\Omega_{PML}}.$$

3. A-priori based $hp$-FEM for eigenvalue problems

It is well known that the accuracy of a finite element approximation of the Helmholtz problem $-\Delta u - \omega^2 u = f$ deteriorates with increasing frequency $\omega$. A major problem is that the discrete frequency of the FE solution is different from the frequency of the exact solution. This effect called pollution has been studied intensively. Particularly, for a uniform mesh size $h$ the asymptotic
error estimates for linear elements [22, Sec. 4.4.3] yield the condition $\omega^2 h < 1$, which for large $\omega$ results in prohibitively expensive meshes. However, the dispersion analysis [22, 23, 24] yields pre-asymptotic estimates of the form $\omega h < 1$, which is a significant improvement. Moreover, it was realized that higher order elements are advantageous to reduce the pollution effect.

A-posteriori estimators are powerful tools when the pollution is negligible, but in the presence of pollution the error in the solution is typically underestimated [25]. For Helmholtz equation with $\omega > 1$ and FE of order $p$ the conditions $p = O(\log(\omega))$ and $\omega h/p = O(1)$ are sufficient for accurate a posteriori error estimation [25]. Recently, error estimates that are explicit in the eigenvalue $\omega$ have also been developed [26]. However, the minimal dimension of a finite element space such that the relative eigenfunction error is below 100% is unknown even in the self-adjoint case with analytic coefficients (see [26, Remark 6.1]).

Our aims are (i) to extend the dispersion analysis in [24] to the case with a complex frequency $\omega$, (ii) to propose an $hp$-strategy for non-self-adjoint eigenvalue problems with piecewise constant coefficients based on the derived conditions in (i) on the finite element space. The a-priori strategy for enriching the finite element space developed in this paper can in principle also be combined with an a-posteriori based strategy such as [27, 28].

3.1. Numerical dispersion for a real frequency $\omega$

The case with a real frequency $\omega$ and constant coefficients has been studied extensively and [22, 23, 24, 29] derived explicit estimates depending only on $\omega$, $h$, and $p$. In this subsection we review those results and consider in the following subsection extensions to complex $\omega$.

In the one dimensional setting, the normalized (wave speed $c = 1$) homogeneous wave equation reads

$$\frac{\partial^2 w}{\partial t^2} - \frac{\partial^2 w}{\partial x^2} = 0. \quad (12)$$

The general solution of the wave equation can be expressed as the superposition

$$w(x,t) = \int_{-\infty}^{\infty} \left[ a(k) e^{i(kx + \omega t)} + b(k) e^{i(kx - \omega t)} \right] \, dk, \quad (13)$$

for some functions $a$ and $b$. The frequency $\omega$ and the wave number $k$ are in this case related by the exact dispersion relation $\omega^2 = k^2$.

We now turn into the numerical computation of (12), where the discrete wave number $k_{hp}$ is a FE approximation to $k$. Let $\{x_j\}, j \in \mathbb{Z}$ be a uniform distribution of points on $\mathbb{R}$, with mesh size $h := x_{j+1} - x_j$, and let $\varphi_j$ be the nodal shape functions of polynomial degree $p$. Then, semi-discrete solutions are written in the form $w_{hp}(x,t) = u_{hp}(x)e^{-i\omega t}$, and at nodal values the FE space representation becomes $u_{hp}(x_j) = \sum_j \xi_j \varphi_j(x_j)$. By analogy with (13), we search for solutions of the form $w_{hp}(x,t) = b_{hp}(k_{hp}) e^{i(k_{hp} x_j - \omega t)}$, which implies $\xi_j = b_{hp} e^{i(k_{hp} x_j - \omega t)}$.

The variational formulation of the problem is then: Find $u_{hp} \in V_{hp} \subset H^1(\mathbb{R})$ such that

$$B_\omega(u_{hp}, v_{hp}) := (u_{hp}, v_{hp}) - \omega^2 (u_{hp}, v_{hp}) = 0, \quad (u, v) := \int_{\mathbb{R}} u \overline{v} \, dx, \quad (14)$$

for all $v_{hp} \in V_{hp}$. The explicit form of $\varphi_j$ in (14) leads to a discrete dispersion relation of the form $\cos(k_{hp} h) = R_p(\omega h)$, where the numerical cosine $R_p(\omega h)$ consists of rational terms involving $\omega$, $h$, and $p$; see [22, 23, 24] for further details. The dispersive error for (12) is defined as $\mathcal{E}^p := R_p(\omega h) - \cos(\omega h)$, from where dispersion analysis refers to studying the convergence of $|\mathcal{E}^p|$ with
respect to $\omega, h$ and $p$. The outcome of the analysis is that $E^p$ is an excellent measurement of the finite element space approximative properties for wave problems as motivated by [22, 23, 24].

We use the following notations: $\kappa = \omega h / 2$, $N_e = \lfloor p / 2 \rfloor$, $N_o = \lfloor (p + 1) / 2 \rfloor$, where $\lfloor x \rfloor$ stands for the integer part of $x$. Ainsworth [24, Sec. 4] proved that when $\omega \in \mathbb{R}$, the function $E^p$ can be written in the form

$$E^p(\omega h) = \frac{\sin \omega h}{\omega h} \left\{ E^p_o \sin^2 \left( \frac{\omega h}{2} \right) + E^p_e \cos^2 \left( \frac{\omega h}{2} \right) \right\} \left\{ 1 + \frac{\sin \omega h}{2 \omega h} (E^p_o - E^p_e) \right\}^{-1},$$

$$E^p_e(\kappa) \frac{\cos^2 (\omega h / 2)}{\omega h} = -Q_{2N_o+3/2}(\kappa) \{ 1 - Q_{2N_o+3/2}(\kappa) \tan \kappa \}^{-1},$$

$$E^p_o(\kappa) \frac{\sin^2 (\omega h / 2)}{\omega h} = -Q_{2N_o+1/2}(\kappa) \{ 1 + Q_{2N_o+1/2}(\kappa) \cot \kappa \}^{-1},$$

with

$$Q_m(\kappa) := \frac{J_m(\kappa)}{Y_m(\kappa)}, \quad m = \text{integer} + \frac{1}{2}.$$  

It was shown [24, Theorem 3.3] that the error $E^p$ for real $\omega$ passes through three phases as the order $p$ is increased: An oscillatory phase, a transition zone, and finally superexponential decay of $E^p$. In the remaining of the section we consider numerical dispersion analysis for $hp$-FEM computations of Helmholtz type of problems with a complex frequency $\omega$. This is of interest since the scattering resonances are complex and the results in this section are the base for the $hp$-FEM strategy proposed in Section 4.2.

3.2. Numerical dispersion for a complex frequency $\omega$

First, we show that the results in [22, 24] can be extended from $\omega \in \mathbb{R}$ to a region in the complex plane. This extension requires that several issues are addressed. Namely, that the expressions can be analytically continued to the complex plane, and the identification of possible branch cuts and poles of the different expressions involved when deriving the estimates in [24]. We rely on the results in [30, 31, 32], where many of the subtleties of working with Bessel functions of complex argument are addressed.

It can be verified that (15) also holds for $\omega \in \mathbb{C}$ with $|\arg \omega| < \pi$. First, by introducing standard sesquilinear forms and following the derivations in [24]. Particularly, equation [24, (4.12)] is reached by using [33, (8.461), (8.465)], which in turn hold for complex arguments. Note that in the case $\omega \in \mathbb{R}$, the subscripts $o, e$ in (15) are reserved for odd, and even, respectively.

3.2.1. Numerical dispersion analysis for small $|\omega h|$

In this subsection, we consider $E^p$ for small $|\omega h|$ and address the case with large $|\omega h|$ in the next section. In the procedure we need the following lemma.

**Lemma 1.** Let $m = n + 1/2$ for $n \in \mathbb{Z}$ and define $Q_m$ as in (16). Then,

$$Q_m(\kappa) = -\frac{1}{2} \left[ \frac{(m - \frac{1}{2})!}{(2m - 1)!} \right] \left( \frac{2m}{2m} \right)^{2m} + \cdots,$$

for all $\kappa \in \mathbb{C}$ with $|\kappa| \ll 1$ and $|\arg \kappa| < \pi$.  


Figure 2: Illustration of Definition 3 and the action of the mapping $g(z)$: Left) Domain $K$, used to characterize the behavior of Bessel functions of complex argument. Right) the domain $D$ is the unit disk.

Proof. We follow the steps in the proof of [24, Lemma A1], which is based on the representation formulas in [33, (8.440)]. Those formulas are under the assumption $|\text{arg}\, \kappa| < \pi$, also valid for complex $\kappa$. Since, in addition $|\kappa| \ll 1$, the representation formulas [33, (8.440), (8.465-1)] hold:

$$J_{\nu}(\kappa) = \left(\frac{\kappa}{2}\right)^{\nu} \sum_{k=0}^{\infty} \frac{(-1)^k}{k! \Gamma(\nu + k + 1)} \left(\frac{\kappa}{2}\right)^{2k},$$

$$Y_{\nu+1/2}(\kappa) = (-1)^{n-1} J_{-\nu-1/2}(\kappa), \quad |\text{arg}\, \kappa| < \pi.$$  \hspace{1cm} (18)

By retaining only the first term in this series and using properties of the $\Gamma$ function, we obtain (17). \hfill\Box

The following theorem extends [24, Thm 3.2] to complex frequencies $\omega$.

**Theorem 2.** Let $p \in \mathbb{N}$, and $|\omega h| \ll 1$. The discrete dispersion relation $E^p$ is then

$$R_p(\omega h) - \cos(\omega h) = \frac{1}{2} \left[ \frac{p!}{(2p)!} \right]^2 \frac{(\omega h)^{2p+1}}{2p + 1} + \mathcal{O}(\omega h)^{2p+4}. \quad (19)$$

Proof. Lemma 1 is stated for complex $\kappa$, with $|\text{arg}\, \kappa| < \pi$, then by plugging (17) into (15), the result follows from the discussion in the proof of [24, Thm 3.2]. \hfill\Box

3.2.2. Numerical dispersion analysis for large $|\omega h|$  

The case $|\kappa| \gg 1$ is of central importance for the paper. To simplify the presentation of this case, we map a particular region of the complex plane on concentric disks.

**Definition 3.** Let $D = \{z : |z| < 1\}$ and denote by $K \subset \mathbb{C}$ the open region enclosed by the parametric curve $w = \pm(\tau \coth \tau - \tau)^{1/2} \pm i(\tau^2 - \tau \tanh \tau)^{1/2}$, $0 < \tau < \tau_0$, where $\tau_0$ is the solution of $\coth \tau = \tau$. For $\delta > 0$, define $S_\delta := \{z : 1 - \delta < |z| < 1 + \delta\}$. Then, we define a continuous bijective mapping $g : \mathbb{C} \to \mathbb{C}$, where the range satisfies $\text{Ran} g|_K = D$ and $g$ maps the set $\{z : \text{dist}(z, \partial K) < \delta\}$ on $S_\delta$. Finally, $g$ is the identity map on $\mathbb{R}$ (see Fig. 2).
Lemma 4. Define $\delta_\nu = (x_2 - x_1)/2$, where $x_1, x_2 \in \mathbb{R}^+$ are the first two real roots of $Y_\nu(x)$. Let $\mathbb{H}^+ := \{ z : 0 < \arg z < \pi, \text{ with } \text{Im} z > \delta_\nu \}$ and $\mathbb{H}^- := \{ z : -\pi < \arg z < 0, \text{ with } \text{Im} z < -\delta_\nu \}$. Then, for $\nu, |z|$ large, and $|\nu g(z/\nu)| > \nu + \nu^{1/3}$ the following approximations hold:

$$Q_\nu(z) \approx \mp i, \quad z \in \mathbb{H}^\pm. \quad (20)$$

Proof. By the conditions stated above, [34, eqs. (9.2.3), (9.2.4)] hold, and $g(z/\nu) \in \mathbb{C} \setminus (D \cup S_\delta)$. From the identities $J_\nu(z) = \frac{1}{2}(H_\nu^{(1)}(z) + H_\nu^{(2)}(z))$, $Y_\nu(z) = \frac{1}{2i}(H_\nu^{(1)}(z) - H_\nu^{(2)}(z))$ and [34, eqs. (9.2.3), (9.2.4)], we obtain the quotient

$$Q_\nu(z) = i \frac{H_\nu^{(1)}(z) + H_\nu^{(2)}(z)}{H_\nu^{(1)}(z) - H_\nu^{(2)}(z)} \approx \mp i \left( \frac{e^{ix}e^{-y}e^{i\theta} + e^{-ix}e^{y}e^{-i\theta}}{e^{ix}e^{-y}e^{i\theta} - e^{-ix}e^{y}e^{-i\theta}} \right), \quad x, y, \in \mathbb{R}, \quad \theta = -\frac{\nu \pi}{2} - \frac{\pi}{4}, \quad (21)$$

where we set $z := x + iy$. Assume that $z \in \mathbb{H}^\pm$, then (21) implies that $Q_\nu(z) \approx \mp i$. \hfill \qedsymbol

The following theorem extends [24, Thm 3.3] to complex frequencies $\omega$.

Theorem 5. Let $p \in \mathbb{N}$, $\omega, \kappa \in \mathbb{C}$, assume that $|\omega h| \gg 1$, and take $\sigma = (2p + 1) \cdot g(\omega h/(2p + 1))$. Then the error $|\mathcal{E}|$ in the discrete dispersion relation passes through three distinct phases as the order $p$ is increased:

i) Non-decaying zone: For $2p + 1 < |\sigma| - o(|\sigma|^{1/3})$, the difference $|\mathcal{E}|$ does not decay as $p$ is increased. For the case with small $|\text{Im} \omega|$, then $|\mathcal{E}|$ oscillates, but does not decay, as $p$ is increased.

ii) Transition zone: For $|\sigma| - o(|\sigma|^{1/3}) < 2p + 1 < |\sigma| + o(|\sigma|^{1/3})$, and $\omega h$ not a pole of (15), the error $|\mathcal{E}|$ decays at rate:

$$|\mathcal{E}| \approx \left| \sin(\omega h) \frac{\text{Ai}(\xi)}{\text{Bi}(\xi)} \right|, \quad \xi = -\left( \frac{2}{p} \right)^{1/3} \frac{\omega h - 2p}{2}, \quad (22)$$

where $\text{Ai}, \text{Bi}$ denote Airy functions.

iii) Superexponential decay: For $2p + 1 > |\sigma| + o(|\sigma|^{1/3})$, $|\mathcal{E}|$ decreases at a superexponential decay rate:

$$|\mathcal{E}| \approx \left| \frac{\sin(\omega h)}{2} f(\sqrt{1 - (\omega h/(2p + 1))^2})^{p+1/2} \right|, \quad (23)$$

where $f : w \to (1 - w)/(1 + w)\exp(2w)$, with $|f(w)| < 1$. In particular, for the case $2p + 1 > |\omega h|e/2$ with $e = \exp(1)$, we have

$$|\mathcal{E}| \approx \left| \frac{\sin(\omega h)}{2} \left[ \frac{\omega h e}{(2p + 1)} \right]^{2p+1} \right|. \quad (24)$$
Proof. In the case $\omega \in \mathbb{R}^+$ is $\sigma = \omega h$ and the theorem was proved in [24, Thm. 3.3]. In the rest of the proof we assume that $\Im \omega \neq 0$ and set $\kappa = \omega h/2$. For $\omega h$ fixed, the error $|\mathcal{E}_p|$ in (15) decays as $|Q_m|$ goes to zero. Then we describe the behavior of $|Q_m(\kappa)|$ in different regions of the complex plane as $p$ is increased.

i) In this regime $|g(\kappa/m)| > 1$, which implies that the point $\kappa$ is in the complement of $K$. Let $\delta_m = (x_2 - x_1)/2$, where $x_1, x_2$ are the first two positive roots of $Y_m(x)$. Then, $Q_m(\kappa)$ is close to a pole on the real line if $-\delta_m < \Im \kappa < \delta_m$. If $|\Im \kappa| < \delta_m$, then by [34, eqs. (9.2.1), (9.2.2)] the function $Q_m$ is oscillatory and dominated by $\Re \kappa$, with an error $e^{\Im \kappa}|O(|\kappa|^{-1})$; Compare with [24, Sec. A.1.1].

For $|\Im \kappa| > \delta_m$, we let $\hat{z} = \kappa/m$, and write $Q_m(m\hat{z})$ in terms of Airy functions. For this, we use the uniform asymptotic expansions [30, (10.20.4),(10.20.5)] with $k = 0$, and neglect the $m^{-5/3}$ term. The trasformation [30, (10.20.3)] can be analytically continued to the complex plane, provided that $\hat{z}$ is located outside $K$. This is true by the assumptions of the theorem. Then, we use [30, (9.6.6),(9.6.8)] to obtain a representation in terms of the Bessel functions $J_{1/3}$, $J_{-1/3}$ of fixed order, which is analogous to [24, (A.7)]. Similarly as in [24, Sect. A.1.1], we use Watson formulas [33, (8.440-1),(8.440-2)] to obtain [24, (A.8)] that holds for complex $\omega$. Finally, the argumentation given in [24, Sect. A.1.1] also holds in the present case. Additionally, Lemma 4 implies that for the current region $|Q_m(\kappa)| \approx 1$, while the order $|\mathcal{E}_p|$ is dominated by $|\sin \kappa|$, which grows exponentially with $|\Im \kappa|$.

ii) As $\kappa$ is not a pole of $Q_m(\kappa)$, we use asymptotic expansions for Bessel functions, which are valid in the transition zone. Particularly, we truncate the series [30, (10.19.8)] with $k = 0$. For $\kappa$ fixed, the resulting formula becomes

\[
Q_m(\kappa) \approx -\frac{\text{Ai}(\xi)}{\text{Bi}(\xi)}, \quad \xi = -\left(\frac{2}{m}\right)^{1/3}(\kappa - m).
\]  

Then for large $p$, we have that $2m \approx 2p$, and with the use of (15) we obtain (22). The same result is obtained by using Olver’s uniform expansions in [30, (10.20.4), (10.20.5)]. As a remark, we mention that even if [24, (A.9)] is a valid linearization of (25), the approximation is quite rough in the complex case because $\min_{m \in \mathbb{Z}+1/2} |\kappa - m| \geq |\Im \kappa|$ may large. However, in the case $\Im \kappa = 0$, the linearization becomes a close approximation to (25) in the transition zone, which implies that $\mathcal{E}_p$ decays algebraically at rate $O(p^{-1/3})$.

iii) In this region $|g(\kappa/m)| < 1$, or equivalently $\kappa/m \in K$. By [31, Sec. 4], it follows that $|Q_m(\kappa)|$ decays. The approximation [24, (A.10)] is justified for complex arguments if the transformations $w = \sqrt{1 - (\kappa/m)^2}$, $z = m(\text{arctanh} w - w)$, and $z = \frac{2}{3}\zeta^{5/2}$ are analytic continuations of its real valued versions. This is easily verified by writing $z \circ w(\kappa)$ explicitly, using the identity $\text{arctanh} w \equiv \log((1 + w)/\sqrt{1 - w^2})$. After a direct calculation we obtain $\sqrt{1 - w^2} = \kappa/m$. Substitution of these into $z/m = (\text{arctanh} w - w)$ results in equation [31, eq. (4.6)], which holds for $z/m \in K$. Finally, by having validated [24, (A.10)], we proceed as in [24] and derive (A.12), (A.13), (A.14), and (A.15). Hence, the results [24, Sec. A.22] hold, which finalizes the proof. Alternatively, (24) is straightforwardly obtained by the use of [30, (10.19.1),(10.19.2)] and (15).

As an illustration of the results in this section, we present in Fig. 3, a dispersion comparison between $\cos(\omega h)$, $R_p(\omega h)$ for $\omega = 20 - 0.5i$ and polynomial order $p = 20$. Followed by a convergence plot showing exponential decay for both real and imaginary parts of the difference $R_p(\omega h) - \cos(\omega h)$. 


Figure 3: In the first two panels we present a comparison of \( \cos(\omega h) \), \( R_p(\omega h) \) for \( \omega \in \mathbb{C} \), by using \( p = 20 \) and show (left) real parts, middle) imaginary parts versus \( h \) in the horizontal axis. In the right panel, we show convergence for \( \text{Re} \mathcal{E}^p \), \( \text{Im} \mathcal{E}^p \) vs. \( p \), with fixed \( h = 2.0 \).

Notice that convergence starts at \( p = 20 \) and \( h = 2 \), where we see in the two first panels that \( h = 2 \) is the largest mesh size where the difference is small.

Dispersion analysis for piecewise constant refractive index:

Consider a problem similar to (14) with \( \omega^2 \) replaced with \( \omega^2 n^2 \), and a refractive index profile \( n \) defined by the constants \( n_j \) for \( x \in I_j \). The problem for \( u \) can then be formulated as the solution of a linear system with matrix entries \( Q_{l,m} \) given by the exponentials \( c_l e^{i\omega x_l} \) defined in \( I_j \). For a piecewise polynomial approximation \( u^{hp} \), we obtain a corresponding matrix \( Q^{hp}(\omega) \) that approximates \( Q(\omega) \). Then, each entry \( \mathcal{E}_{l,m} = |Q_{l,m} - Q^{hp}_{l,m}| \) can be treated similarly as the dispersive error (15), which motivates the use of the dispersion analysis described in Section 3.2.

In the following section we make use of the dispersion analysis revised in the current section, for the FE computation of resonances in one and two dimensions with quadrilateral elements. The natural extension of the discrete dispersion relation to higher dimensions on tensor product meshes is presented in [24, Sec. 2.3]. Particularly, we design a-priori strategies for problems with piecewise constant coefficients, considering each element in our triangulation separately.

4. Discretization, a-priori refinement strategies and solution of the nonlinear eigenvalue problem

In this section we describe the computational details used in order to obtain the approximated resonant pairs as the solution of the nonlinear eigenvalue problems described in section 2. In particular we introduce an initial FE triangulation, which by assumption is conforming and regular. Additionally, we are given a region in \( \mathbb{C} \) where we search for eigenvalues. Then we propose strategies in order to obtain a refined triangulation depending on the permittivity function defined in the computational domain. We motivate the extension to higher dimensions and describe how to obtain the resulting matrix problem in dimensions \( d = 1, 2 \). Finally, we describe our strategy for the solution of the resulting nonlinear eigenvalue problem.
4.1. Discretization with the FE method

The domain $\Omega \subset \mathbb{R}^d$ is covered with a regular and quasi uniform finite element mesh $\mathcal{T}(\Omega_a)$ consisting of elements $\{K_j\}_{j=1}^N$. The mesh is designed such that the permittivity function $\varepsilon(\omega)$ is constant in each $K_j$. Let $h_j$ be the length of the largest diagonal of the non-curved primitive $K_j$ and denote by $h$ the maximum mesh size $h := \max_j h_j$.

In the following, $\mathcal{P}_p$ denotes the space of polynomials on $\mathbb{R}^d$ of degree $\leq p$ and the script $hp := \{h,p\}$ labels the discrete pairs. Furthermore, we assign per element $K_j$ a local polynomial degree $p_j$ satisfying $1 \leq p_j \leq p$. We define the finite element space $S^{hp}(\Omega) := \{u \in H^1(\Omega) : u|_{K_j} \in \mathcal{P}_{p_j}(K_j) \text{ for } K_j \in \mathcal{T}\}$, and $N := \dim(S^{hp}(\Omega))$ [35]. Furthermore, in the case $d = 2$, all our computations are done in the approximated domain $\Omega^{hp}$ by using curvilinear elements following standard procedures [35]. The used FE meshes are shape regular in the sense of [36, Sec. 4.3], and consist of quadrilaterals with curvilinear edges that deviate slightly from their non-curved primitives. Furthermore, we assume that the PML is set up following the discussions in [11, 15], which accounts for large enough $\ell$ and $\sigma_0$ such that the search region is feasible [15]. We assume that the FE space in $\Omega^{PML}$ is good enough and concentrate on the physical region $\Omega_a$.

4.2. A-priori refinement strategies

In the current section, we present two a-priori refinement strategies to be used for the computation of Helmholtz resonances with piecewise constant coefficients. Resonances are then approximated by the eigenvalues of a rational matrix-valued function. The refractive index is approximated by the eigenvalues of a rational matrix-valued function. The refractive index is by computation of Helmholtz resonances with piecewise constant coefficients. Resonances are then approximated in the closed ball $\Lambda := \{ \mu \in \mathbb{C} : |\mu| < 1 \}$ which accounts for large enough $\ell$ and $\sigma_0$ such that the search region is feasible [15]. We assume that the FE space in $\Omega^{PML}$ is good enough and concentrate on the physical region $\Omega_a$.

For the description of our strategy, we use the following definition of an extended mesh.

**Definition 6.** Extended mesh: Let $K_j$ be a one dimensional element of size $h_j := x_2^j - x_1^j$ defined by the nodes in $[x_1^j, x_2^j]$. The extended mesh $\mathcal{M}(K_j)$ is then defined as the partition with points $\hat{x}_l = lh_j$, for all $l \in \mathbb{Z}$.

From this stage we estimate locally the dispersive properties of the finite element space, motivated by the results in Sec. 3.2. Then, refinement strategies can be designed so that for given $k_j$, a finite element space defined over $\mathcal{M}(K_j)$ satisfies the conditions in theorems 2, and 5 for superexponential decay. In this way, we obtain conditions for $p_j$ and $h_j$.

**Definition 7.** For given $k_0$, $h_0 \in \mathbb{R}$, we define the global mesh indicator

$$
\gamma := \left( \frac{k_0 h_0}{2p_0} \right)^{p_0}.
$$
The proposed strategy for enriching the finite element space is based on the goal 8.

**Goal 8.** For given $k_j$ and $\gamma$, find $h_j$ and $p_j$ such that the condition

$$\left( \frac{k_j h_j}{2p_j} \right)^{p_j} \leq \gamma$$

is satisfied in $K_j$.

**Remark 9.** Depending on $k_j$ and $\gamma$, Goal 8 may be unfeasible. Then we say that the target finite element space is unreachable for the given input parameters.

### 4.3. Proposed refinement strategies

The proposed a-priori refinement strategies are based on Goal 8. They guarantee that in each $K_j$, our FE eigenfunctions restricted to $K_j$, satisfy the conditions for superexponential decay of the error on the extended mesh $M(K_j)$.

**Algorithm 1:** A-priori $hp$-FE refinement strategy

**Input:** $p_0$, $K_j$, $h_j$, $n_j$, $\Lambda$, and $\mu \in \Lambda$. Each element is assigned $p_j = p_0$.

1. Compute $k_j$ in each element
2. Set $h_0$ and $k_0$ according to the strategies 4.3.1 or 4.3.2
3. Compute $\gamma$ as defined in (27)
4. Check feasibility of refinement: if $\gamma \geq 1$ then restart with modified input parameters
5. for $j = 1, \ldots, N$ do
6. Check Goal 8 for element $K_j$
7. if Goal 8 not satisfied then refine $h_j$ or $p_j$
8. end
9. Start the assembly of the FE matrices
10. Start the NEP solver with shift $\mu$ and compute the pairs $(u_{hp}^m, \omega_{hp}^m)$

In the following, we present two strategies in order to verify that the condition (28) is satisfied in each $K_j$. The proposed a-priori refinement strategies from the procedure are sketched in Algorithm 1. In the $h$-strategy we assume that $p_j$ is fixed, and we perform a standard $h$-refinement: split the cell $K_j$ in $2^d$ new cells and update $\mathcal{T}(\Omega_a)$ [37]. Similarly, in the $p$-strategy we keep $h_j$ fixed and find a suitable $p_j$.

The initial mesh $\mathcal{T}(\Omega_a)$ is by assumption a conforming triangulation of $\Omega_a$ without ghost nodes; see [37]. Then, a fixed polynomial degree $p_j = p_0 \geq 1$ is assigned to each element $K_j$, $j \in \mathcal{I}_0 := \{j : 1 \leq j \leq N\}$. The refractive index profile $n_j$ is known per element, and the region $\Lambda \subset \mathbb{C}$ containing the shift $\mu$ has been specified. From (26), we assign the constants $k_j$ to each element $K_j$ of mesh size $h_j$.

The parameter $\gamma$ given in (27) determines the state of the initial mesh. In the case $\gamma < 1$, we propose an a-priori refinement strategy, and continue with the steps of Algorithm 1. Otherwise we go back to beginning of Algorithm 1 and ask the user to modify the input parameters.

Below we propose two strategies for achieving Goal 8.
4.3.1. h-strategy (sh-FE)

Let $k_0 = \min_{j \in \mathcal{I}_0} k_j$ and define $\mathcal{I} := \{ j : k_j = k_0 \}$, $h_0 := \min_{j \in \mathcal{I}} h_j$. Then, we perform $h$-refinements of $K_j$, $j \in \mathcal{I}_0$, such that Goal 8 is satisfied. The last statement implies the condition

$$h_j \leq \left( \frac{k_0}{k_j} \right) h_0,$$

from where we proceed to refine all cells until (29) is satisfied. After this, we iteratively refine cells $K_j$ such that each cell has neighboring cells that are at most one level of refinement higher than itself. For this, we allow ghost nodes, and we do not coarsen cells.

4.3.2. p-strategy (hp-FE)

Let $k_0 = \max_{j \in \mathcal{I}_0} k_j$ and define $\mathcal{I} := \{ j : k_j = k_0 \}$, $h_0 := \max_{j \in \mathcal{I}} h_j$. We compute the $p_j$ corresponding to the element $K_j$ such that (28) is satisfied. The last statement implies solving for the zeros $z_i$ of the nonlinear equation

$$F_j(z) := \left( \frac{k_j h_j}{2} \right)^z - \gamma = 0,$$

and choosing the solution $z_i \geq 1$ that minimizes $|p_0 - z_i|$. Finally, we take $p_j := \lceil z_i \rceil$, where $\lceil z_i \rceil$ is the smallest integer greater than or equal to $z_i$.

**Remark 10.** In order to solve (30) we compute the derivative with respect to $z$, and solve by using a scalar Newton-Raphson root finder. We use $z_0 = p_0$ as initial guess and search for solutions in $z_i \in [1, p_0]$. If the only roots are such that $z_i < 1$, then the resulting $z_i$ is not feasible. Possible workarounds are to increase the input parameter $p_0$, or a further uniform $h$-refinement may be needed before starting the strategies.

4.4. Assembly of FE matrices

In this subsection, we consider the assembly of the FE matrices for the 1D problem (7) and the 2D problem (11). Assume that the set of shape functions $\{ \varphi_1, \ldots, \varphi_N \}$ is a basis of the space $S^{hp}(\Omega^{hp})$ defined in section 4.1. Then $u_{hp} \in S^{hp}(\Omega^{hp})$ has the representation

$$u_{hp} = \sum_{j=1}^N \xi_j \varphi_j.$$

4.4.1. Discrete problem in 1D

From (7), with $I^{hp} \subset \mathbb{R}$, we state the corresponding finite element problem: Find $u_{hp} \in S^{hp}(I^{hp}) \setminus \{0\}$ and $\omega_{hp} \in \mathcal{D}$, such that $t_1(\omega_{hp})[u_{hp}, v] = 0$ is satisfied for all $v \in S^{hp}(I^{hp})$.

Similarly, we state the corresponding matrix problem: Find the eigenpairs $(\omega_{hp}, \xi) \in \mathcal{D} \times \mathbb{C}^N$ such that

$$T_1(\omega_{hp})\xi := \left( \sum_{m=0}^{N_r} \left\{ \rho_m(\omega_{hp})A_m - \omega_{hp}^2 \eta_m(\omega_{hp})M_m \right\} - \omega_{hp}\rho_0 E \right)\xi = 0,$$

with finite element matrices

$$A^m_{ij} = \int_{I_m} \varphi_j \varphi_i' \, dx, \quad M^m_{ij} = \int_{I_m} \varphi_j \varphi_i \, dx, \quad E_{ij} = (\varphi_j(-a)\varphi_i(-a) + \varphi_j(a)\varphi_i(a)),$$

for $m = 0, \ldots, N_r$.  

14
4.4.2. Discrete problem in 2D

From (11), with \( \Omega^{hp} \subset \mathbb{R}^2 \), we state the corresponding finite element problem: Find \( u^{hp} \in S^{hp}(\Omega^{hp}) \setminus \{0\} \) and \( \omega^{hp} \in D \), such that \( t_2(\omega^{hp})[u^{hp}, v] = 0 \) is satisfied for all \( v \in S^{hp}(\Omega^{hp}) \). The entries in the finite element matrices become

\[
\begin{align*}
A_{ij}^0 &= (\nabla \varphi_j, \nabla \varphi_i)_{\Omega^{hp}} + (A \nabla \varphi_j, \nabla \varphi_i)_{\Omega^{PML}}, \\
M_{ij}^0 &= (\varphi_j, \varphi_i)_{\Omega^{hp}}, \\
A_{ij}^m &= (\nabla \varphi_j, \nabla \varphi_i)_{\Omega^{hp}}, \\
M_{ij}^m &= (\varphi_j, \varphi_i)_{\Omega^{hp}},
\end{align*}
\]

with \( m = 1, \ldots, N_r \).

The nonlinear matrix eigenvalue problem reads: Find the eigenpairs \( (\omega^{hp}, \xi) \in D \times \mathbb{C}^N \setminus \{0\} \) such that

\[
T_2(\omega^{hp}) \xi := \left( \sum_{m=0}^{N_r} \rho_m(\omega^{hp}) A_m - \omega^{2hp} M_m(\omega^{hp}) \right) \xi = 0.
\]

All numerical experiments have been carried out using the finite element library deal.II [38] with Gauss-Lobatto shape functions [39, Sec. 1.2.3]. For fast assembly and computations with complex numbers the package PETSc [40] is used.

The computational platform used for the executions is Tirant 3, consisting of 336 computing nodes and on each of them two Intel Xeon SandyBridge E5-2670 processors (16 cores each). The processors, running at 2.6 GHz with 32 GB of memory, are interconnected with an Infiniband FDR10 network. All runs are scheduled for at most 4 MPI processes per node.

4.5. Solution of the nonlinear eigenvalue problem

For solving the nonlinear eigenvalue problems we use SLEPc [19] and in particular its NEP module [41]. We provide a target value \( \mu \) and request to compute a few eigenvalues (and corresponding eigenvectors) close to that value. This process is repeated for several values of \( \mu \) in order to cover the region of interest.

The user interface to SLEPc allows the representation of the nonlinear eigenproblem by passing a list of matrices and a list of corresponding scalar nonlinear functions. In our case, the matrix problem to be solved is (35), from where the functions that multiply the matrix coefficients are either polynomial (1 and \( -\omega^2 \)) or rational (\( -\omega^2 \epsilon(\omega) \) and \( 1/\epsilon(\omega) \)). SLEPc provides a simple mechanism to define these functions, either by providing the coefficients of numerator and denominator, or by combining other functions (e.g., additive combination as required in (1)).

In this work, we use SLEPc’s implementation of the NLEIGS method [42], which is based on a two-step process: firstly the nonlinear function is approximated by means of a rational interpolation, and secondly a linearization is built from this rational approximation. Note that since in our case, the function is rational, the rational approximation will be exact if the number of terms is equal to the number of poles plus the degree of the polynomial part. The goal of the linearization step is to obtain a linear eigenvalue problem whose solution is related to the nonlinear one. The dimension of this linear problem is equal to \( d \cdot N \), where \( d \) is the number of terms in the rational approximation and \( N \) is the dimension of the original nonlinear problem. Since this dimension may be quite large, it is important to exploit the structure of the linearization matrices in order to solve the linear problem efficiently (in terms of memory and computational effort). SLEPc’s implementation of NLEIGS represents the Krylov subspace basis in a compact way, \( V = (I_d \otimes U)G \), where vectors of the basis \( U \) have length \( N \) as opposed to length \( d \cdot N \) for vectors of \( V \). The solver builds a Krylov subspace with this basis structure, saving a great deal of memory, and making far
less operations compared to operating with the explicitly formed linearization. In particular, for computing eigenvalues close to the target $\mu$ it is not necessary to factorize a matrix of order $d \cdot N$ but a matrix of order $N$ instead. SLEPc’s implementation also incorporates additional optimizations such as restart and eigenvalue locking. Moreover, all the computation can be done in parallel.

5. Applications to metal-dielectric nanostructures

In this section we study four interesting metal-dielectric configurations, from where numerical approximations to resonances and resonant modes are computed. These configurations are used in Section 6 for comparing the error convergence in standard $h$- and $p$-FE, against the novel strategies presented in Section 4.3. First, geometries with simple symmetries are introduced. This allow us to write exact pairs explicitly, for TM and TE polarizations. Finally, we describe a more demanding problem where this is not possible.

The first two configurations serve as Benchmarking strategies for testing 4.3.1 and 4.3.2 for non-dispersive and piecewise constant material properties. In the last two configurations we are motivated by realistic applications in nano-photonics, where a metal coating is introduced. For these, three different relative permittivity models are used: $\varepsilon_v := 1$ (Vacuum), $\varepsilon_s := 2$ (Silica), and $\varepsilon_{metal}$ (Gold), modeled by a sum of Drude-Lorentz terms (1). For $\varepsilon_{metal}$ we use the data given in table 1 gathered in [43]. This model of Gold has been extensively tested and has validity for $\omega \in [0.5, 6.5]$ eV, where eV denotes electron volt.

5.1. Scaling

In finite precision arithmetic we prefer to work with dimensionless quantities, where we transform from dimensionless variables to physical variables (denoted with $\tilde{}$). We use common physical constants in SI units: $\hbar$ is the scaled Planck’s constant, $c$ is the speed of light in vacuum, and $e$ is the electron charge. In the numerical computations, we use the scaling factors $W = eV/\hbar$ in Hertz and $L = 2\pi c/W$ in meters. Then, we define the dimensionless quantities

$$ x = \frac{\tilde{x}}{L}, \quad \omega = \frac{\tilde{\omega}}{W} \quad \text{satisfying} \quad LW = 2\pi c. \quad (36) $$

The resulting length factor is $L = 1239.842 \text{ nm}$, from where our spectral window becomes numerically equivalent to $eV$ scaling.

\begin{table}[h]
\begin{center}
\begin{tabular}{|c|c|c|c|c|}
\hline
\varepsilon_{\infty} & \omega_p & f_0 & \omega_0 & \gamma_0 \\
\hline
1 & 9.03 & 0.76 & 0 & 0.053 \\
\hline
\end{tabular}
\end{center}
Table 1: Drude Lorentz data for Gold, taken from [43], with time convention $e^{-i\omega t}$.
\end{table}
5.2. Benchmarks in 1D

We focus on the problem described in Section 2.1 for even refractive index profiles. The computational domain is reduced to $I := I_a^+ := (0, a)$, by imposing $u(0) = 0$. This choice allows us to approximate the odd eigenfunctions of (5) and (6). For the derivation of reference solutions, we consider an equivalent coupled problem for $a = 1$.

Let $\{x_j\}_{j=0}^N \subset [0, 1]$ denote nodes with $x_0 = 0$, $x_N = 1$ and introduce the partition consisting of $I_j := (x_{j-1}, x_j)$, $j = 1, 2, \ldots, N$. Assume that the refractive index $n = \sqrt{\epsilon}$ is the constant $n_j$ over $I_j$ and let $u_j$ denote the restriction of $u$ to $I_j$. Furthermore, we assume that $n = 1$, for $x > 1$. Then $(\rho_j, \eta_j) = (1, n_j^2)$ for the TM-case and $(\rho_j, \eta_j) = (1/n_j^2, 1)$ for the TE-case. The coupled problems for the TE/TM-case reads: Find $(u_1, u_2, \ldots, u_N, \omega)$ such that

$$-rac{d}{dx} (\rho_j u_j') - \omega^2 \eta_j u_j = 0, \quad x \in I_j, \quad (37)$$

where $u_1$ and $u_N$ satisfy the boundary conditions

$$u_1(0) = 0, \quad u_N'(1) = i \omega u_N(1), \quad (38)$$

and the solutions of (37) are subject to the compatibility conditions

$$u_j(x_j) = u_{j+1}(x_j), \quad \rho_j u_j'(x_j) = \rho_{j+1} u_{j+1}'(x_j), \quad j = 1, 2, \ldots, N - 1. \quad (39)$$

The general solutions to (37) can be written in the form

$$u_j := A_j e^{in_j \omega x} + B_j e^{-in_j \omega x}, \quad x \in I_j, \quad (40)$$

with the $2N$ unknowns $A_j, B_j$, $j = 1, 2, \cdots, N$. The conditions (38) and (39) imply that the unknowns are solutions of a matrix system

$$Q(\omega) z = 0, \quad z = (A_1, B_1, A_2, B_2, \cdots, A_N, B_N)^T, \quad (41)$$

where the entries corresponding to boundary conditions are placed in the last two rows. If there exist nontrivial solutions to (41), they satisfy det$[Q(\omega_m)] = 0$ for some value $\omega_m$ that corresponds to a resonance of the system.

5.2.1. Slab problem

In this section, we consider the problem (37), (38), and (39) for the case $N = 2$, with $n(x) = n_1$ for $x \in I_1 := (0, 0.5)$, and $n(x) = 1$ for $x \in I_2 := (0.5, 1)$. The corresponding exact resonances for TM polarization are given by

$$e^{2n_1 \omega_m} = -\mu, \quad \omega_m = \frac{(m + 1)\pi - i \text{Log}(\mu)}{2n_1 a}, \quad \mu = \frac{n_1 + 1}{n_1 - 1}, \quad (42)$$

with the corresponding eigenfunctions as in (40), with

$$2 \frac{A_2}{A_1} = (n_1 + 1)e^{i \omega_m(n_1 - 1)} + (n_1 - 1)e^{-i \omega_m(n_1 + 1)}, \quad B_1 = -A_1, \quad B_2 = 0. \quad (43)$$

Similarly, the corresponding exact resonances for TE polarization are given by:

$$e^{2n_1 \omega_m} = \mu, \quad \omega_m = \frac{2m \pi - i \text{Log}(\mu)}{2n_1 a}, \quad \mu = \frac{n_1 + 1}{n_1 - 1}, \quad (44)$$

with the corresponding eigenfunctions as in (40), with

$$2n_1 \frac{A_2}{A_1} = (n_1 + 1)e^{i \omega_m(n_1 - 1)} - (n_1 - 1)e^{-i \omega_m(n_1 + 1)}, \quad B_1 = -A_1, \quad B_2 = 0. \quad (45)$$
TE can be written as degenerated and have algebraic multiplicity where

\[ g(\omega) = 0 \]

The eigenvalues \( \omega_r, \omega_\theta \) in polar coordinates \((r, \theta)\).

### 5.3. Benchmarks in 2D

Accurate Newton reference eigenvalues \( \omega \) for the slab problem described in Section 5.3.2.

| \( m \) | \( j \) | \( \text{Re} \omega_{TM} \) | \( \text{Im} \omega_{TM} \) | \( \text{Re} \omega_{TE} \) | \( \text{Im} \omega_{TE} \) |
|---|---|---|---|---|---|
| 0 | 1 | 1.771 128 241 | −0.040 209 598 | 3.028 519 953 | −0.249 632 742 |
| 1 | 2 | 3.507 165 846 | −0.308 861 246 | 1.276 108 857 | −0.022 849 842 |
| 2 | 3 | 2.637 054 638 | −0.400 052 296 | 1.857 593 240 | −0.103 922 955 |
| 3 | 4 | 3.312 034 818 | −0.666 590 209 | 2.444 174 749 | −0.314 200 015 |
| 4 | 5 | 3.406 691 805 | −0.693 670 033 | 2.506 083 838 | −0.291 213 845 |
| 5 | 6 | 3.525 244 074 | −0.743 331 707 | 2.324 925 787 | −0.200 153 901 |
| 6 | 7 | 3.613 595 702 | −0.818 203 122 | 3.126 303 493 | −0.462 545 189 |
| 7 | 8 | 3.671 987 538 | −0.878 964 710 | 2.510 146 419 | −0.300 384 680 |
| 8 | 9 | 3.720 376 782 | −0.925 532 212 | 5.549 482 036 | −0.741 472 675 |
| 9 | 10 | 3.762 296 208 | −0.963 866 600 | 3.201 508 932 | −0.529 576 832 |

**Table 2: Reference eigenvalues for the single coated disk problem described in Section 5.3.2.**

### 5.2.2. Multiple slab problem

Split the interval \( I := (0, 1) \) in four uniform intervals \( I_j \) of length 1/4 and let \( n := (1, 10, 2, 5)^T \) denote a vector with the refractive indexes \( n_j \). Using this refractive index profile results in an eigenvalue problem that is more demanding for FEM than the slab problem. We compute very accurate Newton reference eigenvalues \( \omega_m \) from \( \det[Q(\omega_m)] = 0 \), with \( Q(\omega) \) given in (41). For simplicity, we only study eigenvalue convergence of this problem for

\[
\begin{align*}
\text{TM} : & \quad \omega_{14} = 10.105 348 365 841 - 0.065 215 027 533 i, \\
\text{TE} : & \quad \omega_{14} = 10.156 176 418 185 - 0.048 229 922 564 i.
\end{align*}
\]

**Equation (46)**

### 5.3. Benchmarks in 2D

The next two problems have radial symmetry centered at the origin, and the solutions expressed in polar coordinates \((r, \theta)\), will be written in terms of Bessel and Hankel functions of integer order \( m \). In this simple case outgoing solutions of (4) satisfy

\[
u = H_m^{(1)}(\omega R) \left( \begin{array}{c} \cos m\theta \\ \sin m\theta \end{array} \right), \quad \text{for } x \in \partial B(0, R), \quad \text{and } m \in \mathbb{Z},
\]

**Equation (47)**

where \( \text{supp}(n - 1) \subset B(0, R) \). In subsections 5.3.1 and 5.3.2, we present solutions satisfying (4) and (47) for specific permittivity profiles.

#### 5.3.1. Single disk problem

Denote by \( u = u_1 \), \( n = n_1 \) the restrictions of \( u, n \) to \( \Omega_1 := B(0, a) \), and set \( n = n_2 = 1 \) elsewhere. The corresponding exact eigenfunctions to (4) and (47) read:

\[
u_1 = N_m J_m(n_1 \omega r) \left( \begin{array}{c} \cos m\theta \\ \sin m\theta \end{array} \right), \quad u_2 = H_m^{(1)}(\omega r) \left( \begin{array}{c} \cos m\theta \\ \sin m\theta \end{array} \right), \quad N_m := \frac{H_m^{(1)}(\omega)}{J_m(n_1 \omega)}.
\]

**Equation (48)**

The eigenvalues \( \omega \) corresponding to \( m = 0 \) are single and those corresponding to \( m > 0 \) are degenerated and have algebraic multiplicity \( \alpha = 2 \). The exact eigenvalue relationship for TM and TE can be written as

\[
J_m(n_1 \omega)H_m^{(1)}(\omega) - g J'_m(n_1 \omega)H_m^{(1)}(\omega) = 0,
\]

**Equation (49)**

where \( g = n_1, \ g = 1/n_1 \) corresponds to the TM polarization and TE polarization, respectively.

18
Figure 4: Convergence plots (Relative errors vs. $N$) for the slab problem 5.2.1 in TM and TE polarizations. The upper horizontal stripe corresponds to classical $h$-FE error convergence for $n_1 = 2, 5, 10$ consecutively. Optimal convergence rates (51) are indicated with solid, dashed, and dotted black lines. The following horizontal stripes correspond to classical $h$-FE and $p$-FE convergence marked with stars, and convergence with the a-priori strategies $sh$-FE (4.3.1), and $hp$-FE (4.3.2) are marked with circles.
5.3.2. Single coated disk problem

In this configuration, we consider a resonator consisting of a dielectric disk with a uniform coating layer. The geometry is described by two concentric circumferences of radii \(0 < R_1 < R_2\), with vacuum as surrounding medium. The inner disk has constant relative permittivity index, and is coated by a layer of gold. We set \(n_1 = \sqrt{\varepsilon_s}\), and \(n_2 := \sqrt{\varepsilon_{\text{metal}}}\) is the value such that \(\text{Im}\{n_2\}\) (absorption coefficient) is positive.

The exact solutions satisfy (4), and (47) with \(R \geq R_2\). The resonance relationship reads

\[
\begin{align*}
\omega_1^m &= g_1 J'_m(\omega R_1) H^{(1)}_m(\omega R_1) - g_2 J'_m(\omega R_1) H^{(1)'}_m(\omega R_1),
\omega_2^m &= g_3 J'_m(\omega R_1) H^{(2)}_m(\omega R_1) - g_4 J'_m(\omega R_1) H^{(2)'}_m(\omega R_1),
\omega_3^m &= g_5 H^{(1)}_m(\omega R_2) H^{(1)'}_m(\omega R_2) - g_6 H^{(1)}_m(\omega R_2) H^{(1)'}_m(\omega R_2),
\omega_4^m &= g_7 H^{(1)}_m(\omega R_2) H^{(2)}_m(\omega R_2) - g_8 H^{(1)}_m(\omega R_2) H^{(2)'}_m(\omega R_2),
F_m(\omega) := (f_1^m f_4^m - f_2^m f_3^m)(\omega) = 0,
\end{align*}
\]

where for TM, \(g := (n_1, n_2, n_2, n_1, 1, n_2, n_2, 1, 1, n_2)\), and for TE, \(g := (n_2, n_1, n_1, n_2, n_2, 1, 1, n_2)\). The parameters used for the computation are \(R_1 = 0.8\), \(R_2 = 1.0\) with scaling factor \(L = 1239.842\) nm.

A complex Newton root finder [44] is then used to compute very accurate approximations of the resonances. For each \(m\) in equation (50), we search numerically the resonances \(\omega_{m,1}, \omega_{m,2}, \ldots\) with machine precision stopping criterion. In Table 2, we list a selection of resonances computed from (50), which are used as a benchmark for studying the proposed hp-FE strategies 4.3.1 and 4.3.2 together with the proposed NEP strategy.

5.4. Coated disk dimer problem

The final configuration consists of two coated disks, each with equal dimension as the one presented in Sec. 5.3.1. The coated disks are surrounded by vacuum, and are separated vertically by a distance \(s = 0.2\). For this problem we compute reference solutions by solving the problem on a very fine mesh.

6. Numerical experiments and results

In this section we perform numerical computations to test the reliability and performance of the proposed solution strategy. Particularly, we present a comparison of classical finite element
Figure 6: Convergence plots (Relative errors vs. N) for TM and TE polarizations: single disk problem 5.3.1 and contrast n_1 = 5. Upper panels correspond to h-FE for p = 2, and bottom panels to p-FE convergence. We mark with circles the a-priori strategies sh-FE and hp-FE, and with stars classical FE refinements. Each vertical strip shows different eigenpairs with j = 1, 2, 4, featuring different angular numbers m = 0, 2, 6.
error convergence against convergence of the a-priori strategies presented in Section 4.3. For the comparison, we define the gain as the percentage of reduction in degrees of freedom compared from using classical FE refinement strategies at a fixed relative error. From a conforming coarse triangulation $T(\Omega)$ with no ghost nodes, the classical $h$ refinement strategy consists in keeping $p$ fixed, and performing consecutive refinements by splitting each quadrilateral in $2^d$ new quadrilaterals. The classical $p$ refinement strategy consists in keeping the number of quadrilaterals constant and increasing $p$ uniformly in each cell.

6.1. Results for non-dispersive problems

The studies are performed on the problems described in Section 5, where expressions for the reference solutions are given for most problems. Furthermore, all given study cases have piecewise analytic coefficients and the domains have no corners. Hence, the expected optimal asymptotic error estimates are:

$$
\|u - u_h\|_l \leq C(\omega) h^{b-l+2} \|u\|_l, \text{ for } h\text{-FE with } l = 0, 1,
$$

$$
\|u - u_h\|_l \leq C(\omega) e^{-\alpha_l N^{1/d}} \|u\|_l, \text{ for } p\text{-FE with } l = 0, 1,
$$

(51)

where we denote by $\|\cdot\|_l$ the standard $H^l(\Omega)$ norm. For the convergence studies, we use the relationship $N \leq ch^{-d}$, $c > 0$ valid for shape regular meshes [36, Sec. 4.3].

In the following sections we discuss the results of the convergence study.

6.1.1. Results for 1D problems

We start by describing general observations resulting from computations on the 1D problems described in Section 5.2. First, we gather results for the single slab problem 5.2.1 in Figure 4.
Figure 8: Resulting polynomial degree distribution \( p_j \) from strategy in section 4.3.2 for shifts \( \mu = 4.162 - 0.2648i \) (left), \( \mu = 2.9 - 0.422i \) (right), and corresponding start values \( p_0 = 7 \) and \( p_0 = 10 \), respectively. In colors we give the computed \( p_j \).

| \( j \) | Re \( \omega_{TM} \) | Im \( \omega_{TM} \) | Re \( \omega_{TE} \) | Im \( \omega_{TE} \) |
|-------|-----------------|-----------------|-----------------|-----------------|
| 1     | 0.391 206 696   | -0.117 682 733  | 1.275 203 310   | -0.017 729 356  |
| 2     | 0.392 635 042   | -0.118 062 545  | 1.407 446 763   | -0.351 566 607  |
| 3     | 0.809 151 314   | -0.171 363 257  | 1.518 834 290   | -0.459 714 126  |
| 4     | 1.775 357 827   | -0.032 801 891  | 1.833 732 651   | -0.071 556 523  |
| 5     | 2.553 994 710   | -0.278 675 516  | 2.122 066 617   | -0.294 491 051  |
| 6     | 2.654 205 934   | -0.403 812 208  | 2.212 801 536   | -0.162 351 126  |
| 7     | 2.889 635 797   | -0.420 955 865  | 2.904 693 880   | -0.427 303 148  |
| 8     | 2.907 250 975   | -0.426 266 060  | 2.905 243 670   | -0.423 913 316  |
| 9     | 3.613 338 577   | -0.811 314 585  | 2.905 244 289   | -0.423 897 437  |
| 10    | 3.668 597 318   | -0.875 944 785  | 3.034 308 619   | -0.251 533 276  |
| 11    | 4.152 064 265   | -0.280 206 808  | 3.104 957 066   | -0.446 726 774  |
| 12    | 4.459 080 468   | -0.244 143 721  | 4.087 887 349   | -0.794 352 424  |
| 13    | 5.565 335 954   | -0.248 309 435  | 4.297 453 212   | -0.260 106 228  |
| 14    | 5.952 378 524   | -0.278 171 454  | 4.491 337 149   | -0.744 646 034  |
| 15    | 6.175 153 470   | -0.890 257 000  | 5.248 059 111   | -0.596 147 639  |
| 16    | 6.636 767 136   | -0.423 048 913  | 5.560 206 560   | -0.251 973 018  |
| 17    | 6.672 655 322   | -0.452 937 916  | 5.809 753 794   | -0.884 187 579  |
| 18    | 7.251 741 950   | -0.616 774 531  | 6.078 619 602   | -0.730 690 710  |
| 19    | 8.155 796 969   | -0.628 012 814  | 7.196 957 492   | -0.638 546 672  |
| 20    | 8.855 927 311   | -0.444 228 537  | 8.961 786 233   | -0.520 547 479  |

Table 3: Reference eigenvalues for the coated disk dimer problem described in Section 5.4.
Figure 9: Left) Spectral window for the coated dimer problem 5.4 with TM polarization. FEM+NLEIGS eigenvalues $\omega_{hp}$ are shown with dots, poles $z$ of $\varepsilon(\omega)$ with squares and its zeros with diamonds. The plasmonic branch points are marked with $\times$ and $+$. Right) In colors we plot $|E_j|$ from the resonant mode corresponding to $\omega_j$ listed in table 3.

Figure 10: Spectral window for the coated dimer problem 5.4 with TE polarization. FEM+NLEIGS eigenvalues $\omega_{hp}$ are shown with dots, poles $z$ of $\varepsilon(\omega)$ with squares and its zeros with diamonds. The plasmonic branch points are marked with $\times$ and $+$. Right) In colors we plot $|E_j|$ from the resonant mode corresponding to $\omega_j$ listed in table 3.
In the upper strip we present classical $h$-FE relative errors corresponding to TM eigenvalues and eigenfunctions measured in $L_2$ and $H^1$ norms. Plots corresponding to $n_1 = 2, 5, 10$ are given for an eigenvalue close to the shift $\mu = 10$. The results indicate that the optimal convergence rates (51) are reached, and the same was observed for the TE polarization. The following horizontal strips depic $H^1$ errors for classical FE and for strategies 4.3.1 and 4.3.2, from where it is observed that both proposed strategies effectively reduce the convergence’s pre-asymptotic phase compared with classical FE refinements. However, the asymptotic rate of convergence remains naturally unaltered. Additionally, optimal convergence rates (51) are reached for both TM and TE polarizations, experiencing the same gain independently from polarization. The second and third horizontal strips in Figure 4 show a gain from $h$-FE of 25%, 44%, and 47%, corresponding to $n_1 = 2, 5,$ and 10 for a relative error around $10^{-3}$. Similarly, the fourth and fifth horizontal strips in Figure 4 show a gain from $p$-FE of 16%, 21%, and 35%, corresponding to $n_1 = 2, 5,$ and 10 for a relative error around $10^{-6}$. The results correspond to $\mu = 10$, but additional numerical computations confirm that similar gains were observed for other shifts. Additionally, we observe from the plots that the gain by using the strategy 4.3.2 increases for higher accuracies. This is expected as the proposed strategies are designed from Theorems 2, and 5. The results confirm that the a-priori strategies 4.3.1 and 4.3.2, achieve convergence with a shorter pre-asymptotic phase than classical FE-methods, with a gain proportional to the refractive index contrast. The results from computations on problem 5.2.2 are gathered in Figure 5, where it becomes evident that the use of the proposed strategies also work well for problems with piecewise constant coefficients. Computations feature a gain of 37% for a relative error of order $10^{-3}$ in $h$-FE, whereas 36% for a relative error of order $10^{-6}$ in $p$-FE.

6.1.2. Results for the single disk problem

In this section the a-priori strategies 4.3.1, and 4.3.2 are tested for configurations in 2D. In particular, the results for the problem 5.3.1, with $n_1 = 5$, are gathered in Figure 6. In this problem, eigenpairs are numbered by using the angular integer $m$ as suggested by (48), and we compute pairs for TM with $\mu = 10.2 - 0.04i$, and for TE with $\mu = 9.85 - 0.04i$. The results for both polarizations are very similar to those discussed in Section 6.1.1 for 1D. Particularly, the eigenvalue error for this problem converges following the optimal rates (51), and both a-priori strategies achieve convergence with a shorter pre asymptotic phase compared to classical FE-methods. Particularly we achieve a gain of 36% for the $h$-strategy with relative error of order $10^{-6}$, whereas up to 17% in the $p$-strategy with relative error of order $10^{-8}$. Furthermore, we see that for the chosen $\mu$, eigenfunctions with different $m$ exhibit the same gain.

Remark 11. For large angular values $m$, eigenfunctions in this problem are expected to exhibit localized oscillations around the boundary of the dielectric disk (juncture with air) that extend to air. These are known as whispering-Gallery-modes (WGM) [45]. It is observed that the strategies 4.3.1, and 4.3.2 underestimate the FE requirements for correct approximation of these modes, as we refine cells according to bulk estimators/goals, and contributions from edges are not considered. However, since we know in advance where to perform mesh refinements it is straightforward to setup a-priori strategies for accurate computation of these modes. From now on, we exclude these type of modes from our discussions.

6.2. Results for dispersive problems

In the remainder of the section we gather results from problems described in sections 5.3.2, and 5.4, which feature dispersive material properties. The positive results from last sections indicate
Table 4: Dimensions and sparsity of the matrix that is factorized during the execution of the NLEIGS solver, for the two test cases of Fig. 11. The number of uniform \( h \) refinements is denoted with \( r \).

| Problem                          | Matrix size | % of nonzeros |
|----------------------------------|-------------|---------------|
| Single coated disk, \( r = 8, p_0 = 2 \) | 4,288,929   | 0.00037       |
| Single coated disk, \( r = 3, p_0 = 10 \) | 173,725     | 0.062         |

that the a-priori strategies 4.3.1, and 4.3.2 applied to non-dispersive problems perform best when there is a high contrast in the refractive index. Similarly, we expect to obtain greater gains when \( |n(\omega)| \) is large. We start by testing the reliability and performance of the NEP solution strategy described in 4.5. Particularly, we check that the strategy can be used to obtain good approximations to the exact resonances even close to the poles and zeros of \( \varepsilon_{\text{metal}}(\omega) \) given in (1). Finally, we consider the error convergence for the problem presented in 5.4, which is computationally more demanding.

6.2.1. Results for the single coated disk problem

In order to test the reliability of the proposed NEP solver strategy, we use the Benchmark presented in Section 5.3.2. From (35), we compute approximations to the resonances given by (50). In Figure 7, we present the result after taking multiple shifts inside a relatively large spectral window, from where we observe an excellent agreement between approximations and exact resonances. We conclude that the proposed a-priori strategies together with SLEPc’s implementation of NLEIGS result in excellent approximations of the exact pairs even close to the poles and zeros of model (1).

Moreover, computations corresponding to TE polarization feature a sequence of resonances accumulating around the so-called plasmonic branch points of the model, which are the values of \( \omega \) such that \( \varepsilon_{\text{metal}} = -1 \), \( \varepsilon_{\text{metal}} = -2 \). For reference, we mark them with \( \times \), and + respectively. From Figure 7 and TE polarization (right), we observe that the approximation \( \omega^{hp} = 0.6288 - 0.6288i \) converged to an eigenvalue of the modified PML problem, which differs considerably from the exact value \( \omega = 0.5569 - 0.6457i \). The reason is that \( \omega^{hp} \) is close to the critical line of the PML [15].

The performance of the solver is evaluated in Fig. 11, where the plots illustrate the strong scaling of the parallel code, that is, how the execution time varies for increasing number of processes with a fixed problem size. Since the problem size is constant, for large number of processes the performance degrades, this because the amount of work assigned to each process is too small. We can see that the run time for 128 processes grows with respect to 64 processes; if the test problems were bigger this performance degradation would occur later for larger number of processes. Still, we cannot expect to scale many more processes since the solver employs a direct linear solver (MUMPS in our case) for one step of the algorithm, which has limited scalability. The figure also shows that the total execution time in the case of higher polynomial degree (right plots) is significantly smaller than for the higher refinement level (left plots). This is due to a much smaller problem size, see Table 4, even though the generated matrices are much less sparse. A shorter time and a higher percentage of nonzero elements also implies a worse scalability, as it can also be seen in the right plots.

6.2.2. Results for the coated dimer problem

Finally, we present results for the coated dimer problem described in Section 5.4, from where the reference values \( \omega_j \) listed in Table 3 were computed from (35) with a very fine discretization. In Figure 8, we show part of the mesh utilized for this problem, and in colors we give the polynomial
distribution $p_j$ per cell resulting by using the $p$-strategy 4.3.2. The distributions shown correspond to $\mu = 4.162 - 0.2648i$ with $p_0 = 7$ (left), and $\mu = 2.9 - 0.422i$ with $p_0 = 10$ (right). We observe that the resulting a-priori strategy assigns lower polynomial degrees to cells with small diameters. As seen from Figure 8, the initial mesh contains a wide range of cell diameters. This property is exploited by the $p$-strategy 4.3.2, because both $h$ and $p$ play a role when satisfying Goal 8.

The resulting a-priori refinement strategy features remarkable gains ranging from 35% to 48% compared to the classical $p$-FE. These gains depend on the selected $\mu$ and on the specific shape of the corresponding eigenfunctions. The error convergence for some of the computed eigenvalues is gathered in Figure 12, where we show convergence for both polarizations and different $\mu$ values. Similarly to the non-dispersive case, the application of the $p$-strategy 4.3.2 to this problem results in shorter pre-asymptotic phase of the error for the computed eigenpairs in both polarizations. Finally, in the left panels of Figures 9 and 10 we present computed eigenvalues from (35) by performing multiple shifts inside a relatively large spectral window. As expected, the location of the resulting eigenvalues resemble those from the single coated disk in Figure 7. Although being more densely populated, the spectral windows exhibit similar features like accumulations to poles, branch points, and similar location of resonances. The Figures 9 and 10 also include color plots for $|E_j(x)|$ corresponding to the $\omega_j$ listed in Table 3, where we have excluded the PML layer. These
plots reveal the rich electromagnetic phenomena described by resonances and resonant modes.

7. Conclusions

We have proposed an $hp$-refinement strategy for approximation of complex scattering resonances in optics. Numerical computations in demanding 1D and 2D cases indicate that the a-priori $hp$-FEM strategy results in a significant reduction of the pre-asymptotic phase in both $h$-FE and $p$-FE. The resulting non-linear matrix eigenvalue problem is solved by SLEPc’s state-of-the-art implementation of the nonlinear eigenvalue solver NLEIGS. This results in fast and highly accurate computations of resonances for metal-dielectric resonators.

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