Efficient periodic band diagram computation using a finite element method, Arnoldi eigensolver and sparse linear system solver.

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Abstract

We present here a Finite Element Method devoted to the simulation of 3D periodic structures of arbitrary geometry. The numerical method based on ARPACK and PARDISO libraries, is discussed with the aim of extracting the eigenmodes of periodical structures and thus establishing their frequency band gaps. Simulation parameters and the computational optimization are the focus. Resolution will be used to characterize EBG (Electromagnetic Band Gap) structures, such as plasma rods and metallic cubes.

Keywords: Finite element method, Eigenvalue problems, Periodic structures, Electromagnetic Band Gap, Sparse matrices.

1. Introduction

The classical plane wave method \cite{1} allows the band diagram of EBG (electromagnetic band gap) structures, like photonic crystal, to be established. These materials are assumed to be periodic, non-dispersive and dielectric ($\epsilon > 0$). However modifications of the plane wave method allow dispersive media \cite{2, 3, 4} to be dealt with. Especially, the plane wave method can also be used to characterize surface mode solutions \cite{5, 6}. However this method, often used in these cases, may be highly difficult to develop when the periodic patterns of the material have an arbitrary shape and permittivity. In this case the Finite Element Method, together with an eigenmode solver, is very efficient. One can, for instance, note the capability of the techniques based on the mesh refinement used by the ANSOFT commercial software HFFS to process it. However, by using this software you cannot control all of the eigenmode resolution parameters \cite{7, 8}. Due to the existence of the spurious modes \cite{9}, the convergence can become very questionable and there is a lack of interpretation and investigation tools to overcome this point\cite{10} and we want to give a detailed explanation about the way to configure the parameters involved in the calculation. The purpose of this study is to propose an effective approach to establish the band-diagram of complex periodical structures. In Section 2 we will explain the theoretical approach of the method, in Section 3 we will explain the way to optimize the Arnoldi algorithm in order to sketch the band-diagram and in Section 4 we will present selected results that illustrate the method.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{example.png}
\caption{Example of a 3d periodic structure with complex shape\cite{11}.}
\end{figure}

2. Scientific basis

Materials are assumed to be non dispersive in this work, that is to say the permittivity is assumed to be non frequency dependent. We are proposing a code that uses a finite element method to solve eigenmode prob-
lems for periodic structures. It will also allow convergence parameters to be controlled at each step of the resolution. For all test function \( \varphi \in H(\text{curl}, \Omega) \) the code is based on the weak formulation [12]:

\[
\int_{\Omega} \frac{\nabla \times \mathbf{E}}{\mu_r} \cdot \nabla \times \hat{\varphi} = k_0^2 \int_{\Omega} \varepsilon_r \mathbf{E} \cdot \hat{\varphi} - \int_{\partial \Omega} (\mathbf{E} \times \frac{\nabla \times \mathbf{E}}{\mu_r}) \cdot \hat{\varphi} \quad (1)
\]

Then we can define the bi-linear forms:

\[
a(a(E, \varphi)) = \int_{\Omega} \frac{\nabla \times E}{\mu_r} \cdot \nabla \times \varphi + \int_{\Omega} (\mathbf{E} \times \frac{\nabla \times \mathbf{E}}{\mu_r}) \cdot \varphi,
\]

\[
m(E, \varphi) = \int_{\Omega} \varepsilon_r \mathbf{E} \cdot \varphi.
\]

Where \( \hat{\varphi} \) denotes the complex conjugate of \( \varphi \).

The unknowns are the electric field \( \mathbf{E} \) and the \( k_0 \) wave number. \( v \) is the outside normal of the interface \( \partial \Omega \). The relative permittivity and permeability are denoted by respectively \( \varepsilon_r \) and \( \mu_r \). We further assume that both relative permeability and relative permittivity are \((3 \times 3)\) positive definite Hermitian tensors. After we mesh the domain we use the Nedelec zero order edge functions[13] on each edge \([i, j]\):

\[
W_{ij} = (\lambda_i \nabla A_j - \lambda_j \nabla A_i) \ast l_{ij}
\]

Where \( \lambda_i \) are the barycentric coordinates of the point \( i \), and \( l_{ij} \) is the length of the edge \([i, j]\). We call \( \mathcal{V}_h \) the subspace generated by these basis functions. Then an approximation of the \( \mathbf{E} \) field can be expressed as a linear combination of these functions:

\[
\mathbf{E} \approx \tilde{\mathbf{E}} = \sum_{i,j} e_{ij} \mathbf{W}_{ij} \text{ where } e_{ij} = \int_{[ij]} \tilde{\mathbf{E}}. \quad (2)
\]

By calling \( \mathbf{X} \) the coordinates of \( \tilde{\mathbf{E}} \) in \( \mathcal{V}_h \) and by applying the Galerkin method, we obtain the linear system:

\[
A \mathbf{X} = k_0^2 M \mathbf{X}, \quad \text{where} \quad A_{pk} = a(W_p, W_k), \quad M_{pk} = m(W_p, W_k), \quad p, k \in [1 \ldots DF], \quad \mathbf{X} = (e_1, \ldots, e_{DF}). \quad (3)
\]

**Remark 2.1.** If we erase the boxed term in (1) then the \( A \) and \( M \) matrices will be positive semi-definite by construction.

\( DF \) are the degrees of freedom of the \( \tilde{\mathbf{E}} \) field, it depends on the boundary conditions. For example, when we impose perfectly electric conducting boundary conditions the integration term on \( \partial \Omega \) disappears and all of the values \( \{e_p, \{p\} \subset \partial \Omega \} \) are equal to zero, the number of unknowns are \( \{e_p, \{p\} \subset \Omega \} \). The purpose is then to compute the eigenmodes first, for metallic cavities (P.E.C boundary conditions) and next for periodic structures by adding PBC (Periodic Boundary Conditions).

After finding solutions for cavities, we have modified matrices initially built to solve problems into periodic structures, by applying the Floquet conditions [1, 14, 15] on \( F_m \) (master) and \( F_f \) (slave) translated by the vector \( \mathbf{T} \):

\[
\mathbf{E}_{F_r} = e^{-j\mathbf{h} \cdot \mathbf{T}} \mathbf{E}_{F_m}, \quad \varphi = k_0 \mathbf{T} \quad (4)
\]

where \( k_0 \) is the incident wave vector, which is an initial condition of the problem, \( \mathbf{E}_{F_r} \) is the field on the slave face and \( \mathbf{E}_{F_m} \) the field on the master face. By definition the band-diagrams are the frequency solutions of Maxwell equations for each given value of \( \varphi \in [0, \pi] \). As the structure is periodic we solve the Maxwell equation on the unit cell of the periodic structure. Let us take an example with a simple bi-periodical structure:

![Example](image)

**Fig.2** In red the basic cell of a 2-dimensional periodic structure. \( T_1 = (a, 0), \quad T_2 = (0, d), \quad k_0 = (k_x, k_y). \)

All the points \( \mathbf{P} \) in the infinite periodic domain have one corresponding point \( \hat{\mathbf{P}} \) in the red domain and if we know the values of the field \( \mathbf{E} \) in this domain, all the solution in the infinite periodic domain can be reproduced using Floquet conditions:

\[
\exists (m, n) \in \mathbb{Z}^2, \quad \hat{\mathbf{P}} = \mathbf{P} + m \times T_1 + n \times T_2 \Rightarrow
\]

\[
\mathbf{E}(\hat{\mathbf{P}}) = \mathbf{E}(\mathbf{P}) e^{-j(k_x T_1 + k_y T_2) \cdot k_0}.
\]

Moreover, the field can be written as a pseudo periodical series [1]:

\[
\mathbf{E}(x, y) = \sum_{n=-\infty}^{\infty} \sum_{m=-\infty}^{\infty} \mathbf{a}_{nm} e^{-j\left(\frac{\pi}{a} x + \frac{\pi}{b} y\right)}
\]

\[
\mathbf{a}_{nm} = \frac{1}{d} \int_0^d \int_0^d \mathbf{E}(x, y) e^{-j\left(\frac{\pi}{a} x + \frac{\pi}{b} y\right)} dx dy
\]
2.1. Consequences on the initial system

In this section, we want to show the transformation of the system (3) induced by the Floquet conditions (4). As we have already seen in equation (4) fields of opposite interfaces are linked by phase shifts and thus, the unknown of the system (3) can be linked as follow :

\[ e_{M_i} = e^{-i\phi}e_{S_i}, \quad \phi = k_iT, \quad k \in [1 \ldots N_M] \]  

(5)

e_{M_i}, \quad k \in [1 \ldots N_M] \] are the unknowns located on \( F_m(\text{master}) \) and \( e_{S_i} \) the corresponding unknowns on \( F_s(\text{slave}) \).

The equation (5) shows that the periodic boundary conditions lead to a reduction in the number of unknowns. As a consequence the approximation subspace will change [16] and we will define the new basis functions :

\[ \tilde{W}_i = W_{I_i}, \quad \tilde{W}_{M_i} = W_{M_i} + e^{-i\phi}W_{S_i} \]

\( i \in [1 \ldots N_I], \quad k \in [1 \ldots N_M] \)

Where \( I_i \) denotes the indices of the internal edges, \( M_k \) denotes the indices of the edges which are located on the master faces and \( S_k \) denotes the indices of the edges which are located on the corresponding slave faces. As a consequence, the variational problem can be written as follows :

\[ \sum_{k=1}^{N_k} e_{M_k} \ast a(W_{M_k}, W_q) + e^{-i\phi} \sum_{k=1}^{N_M} e_{S_k} \ast a(W_{S_k}, W_q) = 0 \]

(\( q = I_i, \quad i \in [1 \ldots N_I] \) or \( q = M_k, \quad k \in [1 \ldots N_M] \))

Remark 2.2. The terms \( a(W_p, \tilde{W}_q) \) will be non-zero terms if the edges \([p]\) and \([q]\) are in the same tetrahedron.

Remark 2.3. The approximation space is equipped with a Hermitian structure :

\[ a(W_p, \tilde{W}_{M_k}) = a(W_p, W_{M_k}) + e^{i\phi}a(W_p, W_{S_k}) \]

More precisely, let us consider a bi-periodical structure, the unknowns are linked by the phase shift relations developed below.

![Fig.3 Phase shift relationships.](image)

The unknowns \( e_{1s}, \) respectively \( e_{2s} \) on the edges on the slave face \( F_{1s} \) (respectively \( F_{2s} \)) are equal to \( e^{-i\phi} \ast e_{1m} \) (respectively \( e^{-i\phi} \ast e_{2m} \)), where \( e_{1m} \) (respectively \( e_{2m} \)) are the unknowns of the field on the face \( F_{1m} \) (respectively \( F_{2m} \)). The phase shifts \( \phi_1 \) and \( \phi_2 \) are respectively equal to \( k_1T_1 \) and \( k_2T_2 \). We see that the edges at the intersection of two slave faces \( F_{1s} \) and \( F_{2s} \) can be correlated with the edges of the intersection of the two master sides \( F_{1m} \) and \( F_{2m} \) and the phase shift will be equal to \( -\phi_1 + \phi_2 \) [15].

The transformed matrices \( \tilde{A} \) and \( \tilde{M} \) by the above method are Hermitian semi-positive definite. Indeed, the above transformations are Hermitian and a matrix \( R \) exists such that [17] :

\[ \tilde{A} = RA^T, \quad \tilde{M} = RM^T \]

Since the matrices \( A \) and \( M \) are positive semi-definite (see remark 2.1) the new build matrices \( \tilde{A} \) and \( \tilde{M} \) will also remain positive semi-definite. If we denote by

\[ INT = \{ I_1, \ldots, I_{N_I}, \quad \text{edge}[I_i] \subset \Omega \}, \]
\[ F_{1m} = \{ M_1, \ldots, M_{N_M}, \quad \text{edge}[M_k] \subset F_{1m} \} \ldots \]

then \( R \) can be written as the bloc matrix :

\[
\begin{bmatrix}
INT & F_{1s} & F_{1s} \cap F_{2s} & F_{2s} & F_{1m} & F_{2m} & F_{1m} \cap F_{2m}
\end{bmatrix}
\]

\[
\begin{bmatrix}
INT & 0 & 0 & 0 & 0 & 0 & 0
F_{1s} & 0 & e^{i\phi_1} & 0 & 0 & i & 0
F_{1s} & 0 & 0 & e^{i(\phi_1+\phi_2)} & 0 & 0 & i
F_{2s} & 0 & 0 & 0 & e^{i\phi_2} & 0 & 0
F_{1m} \cap F_{2m} & 0 & 0 & 0 & 0 & e^{i\phi_2} & 0
\end{bmatrix}
\]

To do this transformation we must construct an appropriate grid :
Remark 2.4. The mesh is structured so that for each member of the master face there is one member of the slave face translated by the constant vector \( T \). Then, if we consider a triangle \( T_m \) on the master face and the corresponding triangle \( T_s \) on the slave face (\( T_m = T_s + T \)), and adding the fact that the tangential component of the basis functions is continuous we have:

\[
\int_{T_m} (\nabla \times (\frac{\nabla W_{qm}}{\mu_r})) W_{pm} = -\int_{T_s} (\nabla \times (\frac{\nabla W_{qs}}{\mu_r})) W_{ps}.
\]

Therefore, the integrals on the master-slave triangles will vanish in pairs, during the assembly of matrix \( A \). For the same reason all of the integrals of the triangles terms will also vanish inside the volume, because the triangles are located at the interface of two tetrahedrons.

Remark 2.5. In comparison with the plane wave method, there is no need to change the permittivity value into the perfect metallic media as has be done in [2]. Indeed the metallic part will be treated with a PEC condition there is no need to mesh this part.

3. Eigen solver strategy

3.1. Storage of the matrices

Throughout the rest of the presentation, we will consider that the finite element matrices are constructed using periodic boundary conditions and we will denote these by \( A \) and \( M \). We can also combine PBC and PEC conditions, but we are not dealing with these possibilities in the examples that we choose to treat. By construction \( A \) and \( M \) are sparse, and we must find an effective way first to store and assemble the matrices. To do this we first have to know the graph of the matrices:

\[
A = \begin{bmatrix}
1 & 2 & 0 & 0 \\
0 & 3 & 9 & 0 \\
0 & 1 & 4 & 0 \\
0 & 0 & 1 & 2
\end{bmatrix}
\]

is a four-by-four matrix with six non-zero elements, thus

\[
\hat{A} = [1 \ 2 \ 3 \ 9 \ 14 \ 12] \\
IA = [1 \ 3 \ 5 \ 7 \ 9] \\
JA = [1 \ 2 \ 3 \ 2 \ 3 \ 4].
\]

3.2. Writing the eigenvalue problem

If the \( A \) matrix is invertible, the linear system can be written as an eigenvalue problem:

\[
AX = k_0^2 MX \Rightarrow A^{-1} MX = \frac{1}{k_0^2} X.
\]  

(6)

Remark 3.1. Usually the central processing unit CPU time for the diagonalization of the matrix \( A^{-1}M \) is proportional to the number of operations needed for the LU factorization of the \( A \) matrix which, is proportional to the cube of its dimension \( N \) and hence to the cube of the number of the plane waves if we apply the plane wave method. However, in our case the matrices are sparse and the number of unknowns are defined by the number of edges in the mesh. Thus, by comparison to
the plane wave method the number of operations and memory used can drastically decrease. Indeed computer storage requirements and the computational complexity of simple array operations are proportional to the number of non-zero elements \( \text{nnz}(A) \) [19]. Looking at the LU factorization for sparse matrices which uses a zero order reduction algorithm [20], we can deduce that the number of operations for the diagonalization of a sparse matrix should be proportional to \( N \times \text{nnz}(L + U) \approx N \times \text{nnz}(A) \).

The algorithm we choose to use will give the largest eigenvalues and these eigenvalue will be equal to \( \frac{1}{k_0^2} \), so we will get the first few smallest values of \( k_0 \). However, the transformation shown in equation (6) is not suitable because the matrix \( A \) is not invertible. Indeed, thanks to the discrete Helmholtz decomposition [21], there exists \( U \in (H^1(\Omega))^3 \) derived from a vector potential \( V \in H(\text{curl}, \Omega) \) and a scalar potential \( \phi \in H^1(\Omega) \) such as:

\[ \tilde{E} = U + \nabla \phi, \text{ such as } \nabla \cdot U = 0 \text{ and } U = \nabla \times V. \]

The kernel of \( A \) then contains the fields \( \tilde{E} \) (defined equation (2)) which have a zero rotational component (i.e \( U \equiv 0 \)). More precisely, the solutions of the equation (3) may be classified into 3 groups [9]:

\[
\begin{align*}
\text{Group 1} & : k_0^2 \neq 0 \text{ and } \nabla \cdot [ \epsilon_r ] \tilde{E} = 0 \\
\text{Group 2} & : k_0^2 = 0 \text{ and } \nabla \cdot [ \epsilon_r ] \tilde{E} = 0 \\
\text{Group 3} & : k_0^2 = 0 \text{ and } \nabla \cdot [ \epsilon_r ] \tilde{E} \neq 0
\end{align*}
\]

The solutions included in Group 3 are called spurious modes and we want to avoid them. These solutions can be written as a potential-like function and they have a zero curl. When we solve cavity problems, the spurious modes can be eliminated by adding a zero divergence constraint into the Lanczos algorithm [22]. When we deal with periodical boundary conditions, the technique used in [22] is no longer feasible with zero order Nedelec basis functions because of the constraint on the meshes [23]. In our problems, we will eliminate all of the zero solutions, that is to say, we will keep only the solutions included in Group 1. As the kernel of the matrix \( A \) is not equal to zero, the initial eigenvalue problem (6) must be transformed. The transformation we have decided to apply is the shift and invert transformation, which consist in computing the eigenvalues of the operator \((A - \sigma M)^{-1}M\). If we denote by \( \mu \) one eigenvalue of the operator \((A - \sigma M)^{-1}M\), we have:

\[
(A - \sigma M)^{-1}M = \mu X \text{ with } \mu = \frac{1}{k_0^2 - \sigma}
\]

Then, if we seek solutions for the problem \( AX = k_0^2 M X \) we must look for the greatest values of the problem \( OP \ast X = \mu X \) and the values obtained will be close to the shift \( \sigma \), given \( OP = (A - \sigma M)^{-1}M \).

To do this, we use an improved Arnoldi algorithm proposed by the ARPACK library [24] on operator \( OP \). Arnoldi algorithm consists in orthonormalizing the Krilov space \((\text{span} \{ v_0, OPv_0, OP^2v_0, \ldots, OP^{n-1}v_0 \})\) and is as follows:

\[
\begin{align*}
\text{w} & = OP \ast v_j \\
& \quad \text{for } i = 1 \text{ to } j\text{ do} \\
& \quad h_{ij} = (w, v_i) \\
& \quad w = w - h_{ij} v_i \\
& \quad \text{end do} \\
& \quad \text{h}_{j+1, j} = \| w \| \\
\end{align*}
\]

At each step we have to compute the value of \( OPv_j \). In other words we must solve the linear system:

\[
(A - \sigma M)X = Mv_j
\]

(9)

First we have to compute the sparse matrix-vector product \( Mv_j \) which will be stored in the vector \( Y \) (the notations are the same as in section 3.1):

\[
\begin{align*}
Y & = 0 \\
\text{for } i = 1 \text{ to } n \text{ do} \\
Y(i) & = Y(i) + \tilde{M}(j) \ast XJM(j) \\
\text{END DO} \\
\text{END DO}
\end{align*}
\]

Here we can see that the cost of operation to perform a matrix vector product is very cheap. Then we have to solve the linear system (9). To do this we choose to use a direct sparse solver PARDISO [25] which uses zero order reducing algorithms to first perform the LU factorization to the matrix \((A - \sigma M)\) and then solve the system. As we have triangular matrices (lower and upper) the system can be solved at each step directly using forward and backward resolution.

**Remark 3.2.** In general the structure of the graph of sparse matrices influence the computational time of the LU factorization performed by zero order reducing algorithms. Even if we can optimize these types of algorithms based on profiles of matrices, the graph structure of the matrix \( A \) will influence the computational time of solving the system (9). As we have already noticed in section 3.1, the graph structure is directly related to the structure of the mesh. We can therefore conclude that the structure of the mesh affects the computation time of the resolution of the system (9).
3.3. Properties of the construction

After the construction of the first Arnoldi vectors, we have the following properties:

\[ OPV_j = \sum_{i=1}^{j+1} h_{ij} V_i. \]

If denote by \( V^p = (v_1 \ldots v_{\bar{p}}) \), we have \( OP \ast V^p = V^p_{p+1} H_{p+1} \) with:

\[
H_{p+1} = \begin{pmatrix}
   h_{1,1} & h_{1,2} & \cdots & h_{1,j} & h_{1,\rho} \\
   h_{2,1} & h_{2,2} & \cdots & h_{2,j} & h_{2,\rho} \\
   0 & h_{3,1} & \cdots & h_{3,j} & h_{3,\rho} \\
   \vdots & \vdots & \ddots & \vdots & \vdots \\
   \vdots & \vdots & \ddots & \vdots & \vdots \\
   0 & \cdots & \cdots & h_{j+1,j} & h_{j+1,\rho} \\
   \vdots & \vdots & \ddots & \vdots & \vdots \\
   0 & \cdots & \cdots & 0 & h_{p+1,\rho}
\end{pmatrix}
\]

Then let the upper Hessenberg matrix \( H^p \) be the matrix \( H_{p+1} \) without the \( \bar{p} + 1 \) row. We have the equalities:

\[ OP \ast V^p = V^p H^p + h_{p+1,\rho} V^p_{p+1} \otimes e^p \]  \hspace{1cm} (10)

This can also be written as:

\[ (OP - h_{p+1,\rho} V^p_{p+1} \otimes v^p) V^p = V^p H^p. \]

If we denote by \( Z^p \) one eigenvector of the matrix \( H^p \), \( V^p Z^p \) is an eigenvector of the operator \( OP \) and we have the following approximation:

\[ \| OP \ast V^p Z^p - \lambda V^p Z^p \| \leq \| OP \ast V^p \| \| V^p H^p Z^p \| \leq |h_{p+1,\rho}| \| V^p_{p+1} \| \| Z^p \| \]

Let \( \tilde{\sigma} \) be a fixed real positive quantity. It is considered that eigenvalue is acceptable if

\[ |h_{p+1,\rho}| \| V^p_{p+1} \| \| Z^p \| \leq \tilde{\sigma}. \]  \hspace{1cm} (11)

Afterwards the Implicit Restarted Arnoldi Algorithm also called IRAM can be written as follows [26]:

Start: Build a length \( \bar{p} \) Arnoldi factorization \( OP \ast V^p = V^p H^p + h_{p+1,\rho} V^p_{p+1} \otimes e_p \) with a starting vector \( v_1 \).

1. Compute the eigenvalues \( \{ \lambda_j : j = 1, 2, \ldots, \bar{k} \} \) by using classical QR algorithms [27] combined with Givens rotations [28, 29] applied to the matrix \( H^p \).
2. If the first \( \bar{k} \) eigenvalues verify the criterion (11), then stop the algorithm.
3. Else Perform \( \bar{p} - \bar{k} = m \) steps of the QR algorithm on the matrix \( H^p \) with the unwanted eigenvalues \( \{ \lambda_j : j = \bar{k} + 1, \bar{k} + 2, \ldots, m \} \) as shifts [30] to obtain \( H^p Q^p = Q^p H^p \).
4. Restart: Post multiply the length \( \bar{p} \) Arnoldi factorization with the matrix \( Q^p \) consisting of the leading \( \bar{k} \) columns of \( Q^p \) to obtain the length \( \bar{k} \) Arnoldi factorization \( OP \ast V^p Q^k = V^p Q^k H^p \) + \( \sum_{j} \bar{k} \ast e^p \), where \( \bar{k} \) is the leading principal sub-matrix of order \( \bar{k} \) for \( H^p \). Set \( V^k \leftarrow V^p Q^k \).
5. Extend the length \( \bar{k} \) Arnoldi factorization to a length \( \bar{p} \) factorization and return to 1.

3.4. Solution set up

Let us make a quick summary about the use of the solver. The input parameters are:

| Two sparse and Hermitian finite element matrices. | \( \tilde{\Lambda} \) and \( \tilde{M} \) |
|------|------|
| Phase shift between master slave faces | \( \phi = k_i T \) |
| Number of wanted eigenvalues (Also called Ritz values) | \( \bar{k} \) |
| Number of Arnoldi vectors | \( \bar{p} \) |
| Tolerance criteria | \( \tilde{\sigma} \) |

A good choice of the initial shift is given by:

\[
\sigma = 3 \pi^2 \min \left\{ \frac{1}{L^2}, \frac{1}{h^2} \right\} \sum_{V_i \in \text{eI}} \frac{1}{V_i} V_i \]

Where \( L \) correspond to the length, \( l \) the width and \( h \) to the height of the domain. \( \epsilon_r, \mu_r \) are the relative permittivity and permeability of the different sub-domains and \( V_i \) is their respective volume. This choice is motivated by the fact that the value of the first eigenmodes in an homogeneous rectangular cavity filled with a dielectric
\((\epsilon_r, \mu_r)\) is equal to [31]:

\[
\left( k_0^2 \right)_{\text{map}} = \frac{\pi^2}{\epsilon_r \mu_r} \left( \frac{p^2}{L^2} + \frac{m^2}{T^2} + \frac{n^2}{h^2} \right).
\]

The value of the shift \(\sigma\) approximately corresponds to the first analytical eigenmode in a rectangular metallic cavity of the same size. The initial value given by the relation (12) is only an estimation used as a point of departure for the solver. This is the reason we let the user the possibility to change this initial value. For example, when the periodic cell contains several resonant elements, the shift can be adjusted to the wave number proportional with the wavelength corresponding to the size of one of these elements. Moreover, it is easy to understand that the parameter \(\sigma\) depends on the number of wanted eigenvalues \(\tilde{k}\). More precisely, if we increase the value of \(\tilde{k}\), we can choose \(\sigma\) further from to the first wanted eigenvalues. Moreover, since the parameters are linked-together they do not all need to be filled. Indeed the numerical accuracy of the result will depend on the convergence criteria \(\text{tol}\), the number of Arnoldi vectors \(\tilde{p}\) and the number of wanted eigenvalues \(\tilde{k}\). Due to the structure of the algorithm, we can easily understand that \(\tilde{p}\) must be greater than \(\tilde{k}\) and \(\tilde{p}\) must be large enough to avoid too many restarts of the IRAM [26] algorithm. This is the reason we choose to link these two parameters as follows:

\[
\text{Remark 3.3. For each eigenmode solution } (k_0^2, X), \text{ we introduce the residue } \epsilon \text{ which will allow to eliminate null eigenvalues according to the criterion :}
\]

\[
\text{if } \epsilon = \frac{\|AX - k_0^2MX\|}{\|k_0^2\|} \leq \text{tol} \quad \text{(13)}
\]

then \((k_0^2, X)\) is an acceptable solution.

It is well known that the number of unknowns will influence the physical accuracy of the result. Moreover, some part of the mesh domain has to contain more edges than others in order to compute values that correspond to the physical solutions. In order to know exactly which part we have to mesh thinner we can iterate on the mesh and compare the local variations of the solutions as HFFS do [7]. By doing this process we can observe than the thinner meshes have to be located in areas with highly diffracting materials or areas "narrow" where geometry varies abruptly. But this process is very time expensive, so we just need to get in a first step the optimized mesh.

3.5. Sort algorithm

We present here an algorithm used to remove the zero solutions by changing the value of the shift and using the criterion (13) during the calculation:

- Thanks to the Arnoldi algorithm, we compute the eigenvalues of the operator \((A - \sigma M)^{-1} M\) where the initial shift \(\sigma\) is defined by the relation (12).

- We will obtain \(\tilde{k}\) Ritz values for each given value of \(k_0\).

- \(n_s\) of these solutions fulfill the convergence criteria (13). These \(n_s\) solutions are stored in the array called \(\text{Tab}[\].

- If \(n_s < 2 \cdot \tilde{k}/3\), the value of the shift \(\sigma\) increased , the solver is restarted and the values are not retained. For a fixed \(\tilde{k}\) we change the value of \(\sigma\) (called \(\sigma_{\text{new}}\)) compared with the old value \(\sigma_{\text{old}}\):

\[
\sigma_{\text{new}} = \frac{1}{4} \max(\text{Tab}[\] + \frac{3}{4}\sigma_{\text{old}}
\]

- Else If \(n_s = \tilde{k}\) the value of the shift \(\sigma\) decreased , the solver is restarted and the values are not retained. For a fixed \(\tilde{k}\) we change the value of \(\sigma\) (called \(\sigma_{\text{new}}\)) compared with the old value \(\sigma_{\text{old}}\):

\[
\sigma_{\text{new}} = \frac{1}{4} \min(\text{Tab}[\] + \frac{3}{4}\sigma_{\text{old}}
\]
• Else the values are retained and we change the value of \( \sigma \) (called \( \sigma_{\text{new}} \)) compared with the old value \( \sigma_{\text{old}} \):

\[
\sigma_{\text{new}} = n_s - 1 \sigma_{\text{old}} + \frac{\overline{Tab}}{n_s},
\]

where \( \overline{Tab} \) means the average value of \( Tab \).

Finally, for each value of \( k_i \), we will print all eigenvalues \( (k_{i0}, X) \) that fulfill the convergence criteria (13) in a band diagram. We can also print the field created for each Ritz value. If \( \sigma \) is chosen too large, all solutions will verify the criteria (13), i.e. \( n_s \) is equal to \( \bar{k} \) and the algorithm is restarted until \( n_s < \bar{k} \). Thus the choice of the initial value of shift \( \sigma \) will play an important role in the computation time.

4. Results

4.1. Metallic Cubes

We consider perfectly conducting cubes left in air. The cell size is the original cube height \( D_x = D_y = D_z = 1 \text{m} \), and the length of the side of the cube \( w = 0.5 \text{m} \). The characteristics \( \epsilon, \mu \) are those of the air \( \epsilon_1 = \mu_1 = 1 \). The value of the shift \( \sigma \) determined by (12) is equal to 5 rad/m.

For our simulations, we first vary the phase shift in the \( Ox \) direction from 0 to \( \pi \) and the phase shifts in the other directions are set to zero. Then, we fix the phase shift in the \( Ox \) direction to \( \pi \), that in the \( Oz \) direction to 0 and that in the \( Oy \) direction varies from 0 to \( \pi \) (range \([\pi, 2\pi]\) in the diagram). Then, the phase shifts \( Ox \) and \( Oy \) are fixed at \( \pi \) and the \( Oz \) phase shift varies from 0 to \( \pi \) (range \([2\pi, 3\pi]\) in the diagram). The results are expressed in radians per metre and the phase is given in radians. On the metallic parts, the tangent field is null.

In Fig.8 the method used in the Article [14] is also a finite element method, but the algorithm used to compute the eigenmodes is not specified. In Fig.8 the number of unknowns is equal to 10344 and in Fig.9 9599. This explains the very small difference between the two diagrams. To conclude this example, we can say that the results are validated and that it is possible to extend the method to non-canonical geometries.

Remark 4.1. The FDTD[32, 33] method can also treat this kind of problem but if we change metallic cubes to 3D complex structures (cylinders or pyramids for example); since the meshed area is necessary cubic and the geometry must be meshed uniform way and small enough to ensure the stability of the numerical scheme [33], computation time and memory space required can drastically increased. Our method accepts all of the kinds of meshes, unless the master slave relations are respected on the master slave faces. Thus we have less restrictions on the geometry.
4.2. Infinite dielectric rods

**Basic cell.** Let us consider a square network of infinite dielectric rods in the $Oz$ direction. The entries on the axes are the components of the plane wave excitation.

![Square network of infinite dielectric rods](image1)

Fig. 10 Square network of infinite dielectric rods $a=7$mm, radius=2mm, $\epsilon_r = 9.4$.

![Phase shift](image2)

Fig. 11 We vary the phase shift in the basic cell. The phase shift in the $Oz$ direction is set to 0. The shift $\sigma$ defined by (12) is equal to 21 Giga Hertz.

The geometry is invariant in the $OZ$ direction, then we set $k_{iz} = 0$ and the height of the meshed area is chosen thinner than the length and the width:

![Meshed area](image3)

Fig. 12 Meshed area of the basic cell with a thickness=1mm.

As we can see, the plane wave diagram can be accurately reprinted with our method. To treat this case, it is also possible to use the finite element method for two-dimensional doubly periodic structures, as it is developed in [14]. Due to the geometry, the solution can easily be computed with the plane wave method using Bessels functions [1].

**Remark 4.2.** We can deal with this problem by using HFFS, but in the latest version the solver will not give the solutions under the minimum chosen frequency [8]. This can be problematic if the first solutions are very near to zero, as we can see in this case because if we set this frequency to a small value the results will be corrupted [7]. In addition, if this frequency is set to a value that is too large, we will not get the first eigenmodes.
4.3. Super-cells

Periodic patterns which will have more than one basic cell are called super cells. The basic cell is now defined by $a = 14 \text{mm}$, radius $r = 0.18a$ and $\epsilon_r = 11.56$. We want to study the propagation in the $Ox, Oy$ plane (Fig.16, Fig.20), so the $k_x$ component is fixed to zero and the thickness of the cells is equal to 1mm. We vary the phase shift $k_y = \phi_y/\text{dy}$ for multiple values of $k_{xx} = \phi_x/\text{dx}$. In other words, we compute the projected band diagram [6] in the $Oy$ direction.

In Fig.15, the straight red line represents the light line defined by the equation $\omega = c|k|$. We call guided modes that are able to propagate in the air but not into the meta-material. These appear when we introduce a linear defect into the material; the existence of these modes corresponds to the green zone. Modes that are evanescent in the air and in the meta-material are called surfaces modes. These correspond to the yellow zone and appear when we change the size of the last elements of the periodical structures [6]. To trap these modes, we establish the band diagrams of super-cells [6]. The $Oy$ direction corresponds to the direction of the defect introduced in the super-cells.

![Fig.15](image)

**Remark 4.3.** The shift $\sigma$ defined by (12) will be too small because the dimensions of the super cell are much larger than the dimensions of the basic cell. Then, we choose a shift included in the band gap that we had already found by studying the basic cell.

The corresponding band-diagram for the super-cell Fig.16 is:

![Fig.16](image)

In Fig.17 each point $(\phi_x, \phi_y)$ will give 12 over 15 eigenvalues which will verify the criteria (13).

Here we can observe the values of the $\text{Re}(E_z)$ and $|E_z|$ guided fields on the plane $z=0.0005$. The frequency of the mode is 8.58 Giga Hertz:

![Fig.17](image)

![Fig.18](image)

![Fig.19](image)
4.3.1. Surface defect

We now will introduce a defect by modifying the last row of the super-cell:

![Fig.20 Modification of the last rods to make surface modes appear. The radius of the surface rods is equal to 0.009\(a\).](image)

Remark 4.4. In [6, 5] the plane wave method is used to characterize surface modes when the network is defined by square rods or circular rods. This emphasizes the fact that the last rod must be different from the rods of the network, in the example the last rod is cut down its middle.

The corresponding band-diagram for the Fig.20 is:

![Fig.21 Projected band diagram for a super cell with modification of the surface rods in the Oy direction. \(\phi_x \in [0, \pi, \pi/2, 3 \pi/4]\) and \(\phi_y \in [0, \pi/8, \pi/4, \pi/2, \pi]\).](image)

Remark 4.5. By modulating the value of the shift \(\sigma\), we can directly seek solutions that are located in the red and yellow part and then there is no need to consider a lot of wanted eigenvalues \(\vec{k}\). For example, if we only seek surface mode solutions we can take \(\vec{k} = 2\) and \((\sigma, \phi_y)\) located in the yellow zone in Fig. 15.

5. Summary of the computational performances

We now make a summary for each case of the computational performances treated on our Intel (R) Xeon 64 bits 2.80 GHZ machine. We specify the CPU time, the total number of edges in the mesh file, the RAM used during the process and the number of points in the band-diagram. The number of wanted eigenvalues \(\vec{k}\) is always equal to 15 and the tolerance criteria \(\text{tol}\) is always equal to \(10^{-7}\).
6. Conclusion and perspective

We have presented the theoretical aspect of the eigenmode solver for periodic non-dispersive structures first. Then, we have presented the way to combine different libraries, such as ARPACK and PARDISO, to optimize the Arnoldi algorithm and finally, we have presented the results on the original periodic structures. We are now trying to study a dispersive structure by adapting a non-linear plane wave technique [3] to our model.

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