Research Article

Integral Equation Analysis of EM Scattering from Multilayered Metallic Photonic Crystal Accelerated with Adaptive Cross Approximation

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A space-domain integral equation method accelerated with adaptive cross approximation (ACA) is presented for the fast and accurate analysis of electromagnetic (EM) scattering from multilayered metallic photonic crystal (MPC). The method directly solves for the electric field in order to easily enable the periodic boundary condition (PBC) in the spatial domain. The ACA is a purely algebraic method allowing the compression of fully populated matrices; hence, its formulation and implementation are independent of integral equation kernel (Green’s function). Therefore, the ACA is very well suited for accelerating integral equation analysis of periodic structure with the integral kernel of the periodic Green’s function (PGF). The computation of the spatial-domain periodic Green’s function (PGF) is accelerated by the modified Ewald transformation, such that the multilayered periodic structure can be analyzed efficiently and accurately. An effective interpolation method is also proposed to fast compute the periodic Green’s function, which can greatly reduce the time of matrix filling. Numerical examples show that the proposed method can greatly save the frequency sweep time for multilayered periodic structure.

1. Introduction

Photonic crystals [1] are periodic structures of great interest for their applications both in the microwave region and in the optical range. The main feature is the presence of frequency bands wherein the waves are highly attenuated and cannot propagate. It results from the removal of degeneracies of the free-photon states at the Bragg planes provoked by the periodicity, which produces forbidden frequency gaps so-called photonic band gaps (PBGs). This property is exploited in the electromagnetic and optical applications.

Many numerical methods for computing such band structures have been adapted from solid-state physics. They include, but are not restricted to, plane-wave expansions [2], the Korringa-Kohn-Rostoker method [3], and shell methodologies [4]. Over the past two decades, finite difference time domain (FDTD) [5] has been widely used in computational optics and photonics. Recent developments include specialized versions of the finite element method (FEM) [6] and the flexible local approximation method [7]. FDTD and FEM can cope with arbitrary complicated shapes and materials. However, the radiation into unbounded regions requires either absorbing boundary conditions or perfectly matched layers, and special measures need to be taken into account when employing these conditions to the scattered field formulation. For open-region problems, integral equation/method of moments (IE/MoM) has a great advantage.

The ACA is a purely algebraic method, which is independent of the kernel. The ACA was originally introduced in 2000 by Bebendorf [8] and since then has been successfully applied for integral formulations involved electromagnetic scattering [9, 10]. The integral kernel is a free space Green’s function. In addition, it is noted that most photonic crystal
applications deal with two-dimensional (2D) structures that are invariant along a longitudinal axis and periodic in the transverse plane [11]. The manufacture of a 2D photonic crystal structure is easier than that of three-dimensional (3D) one [12].

In this paper, a space-domain integral equation accelerated with ACA is presented for the fast and accurate analysis of electromagnetic scattering from 2D and 3D multilayered periodic structures. The ACA is first introduced for surface integral equation with the kernel of periodic Green’s function (PGF). The ACA can be applied to compress impedance matrix blocks and accelerate matrix-vector multiplication (MVP) for reducing computational complexity. A modified Ewald transformation is proposed to efficiently compute the PGFs. Compared with traditional Ewald method, it has less computational cost and can eliminate the imbalance of two Ewald sequences for multilayered periodic structure. An efficient interpolation method is also proposed to fast compute the PGFs and then accelerate impedance matrix filling. Our proposed method can effectively reduce the frequency sweep time for multilayered periodic structure involving a lot of unknowns.

2. Formulation

Figure 1 shows the multilayered double periodic structure with identical metallic objects for arbitrary shape periodically repeated in the xy-plane. $\mathbf{a}_1$ and $\mathbf{a}_2$ are the primitive lattice vectors. $A = |\mathbf{a}_1 \times \mathbf{a}_2|$ is the cross-sectional area of the unit cell. $\mathbf{E} = (\hat{\theta} \cos \alpha + \hat{\varphi} \sin \alpha)E_0 \exp(-j \beta \mathbf{k} \cdot \mathbf{r})$ is the incident electric field. $\alpha$ is the polarization angle. The wave vector is $\mathbf{k} = \beta \mathbf{k}$ and phase shift $\mathbf{k}_{100} = \mathbf{k}_1$. The array is assumed to be periodic in the xy-plane and the cell of the array is obtained by shifting the cell through the relation $\mathbf{p}_{mn} = m \cdot \mathbf{a}_1 + n \cdot \mathbf{a}_2$. $\rho_{mn}$ is defined as the translation vector of the lattice.

2.1. Integral Equation. In this paper, we consider the metallic photonic crystal. Let S be the surface of a metallic object. By enforcing the boundary conditions that the total tangential electric field is zero on the perfect electric conductor (PEC) surface S, the electric field integral equation (EFIE) is given as [13]

$$ j \omega \mu_0 \left[ \int \mathbf{J}_s(\mathbf{r}') \frac{1}{\mathbf{p}^2} \mathbf{e}^{j \mathbf{p} \cdot \mathbf{r}'} \cdot \mathbf{G}_p(\mathbf{r}, \mathbf{r}') \, ds' \right] = E_0^{inc}, $$

(1)

where $G_p$ is a doubly periodic Green’s function. The Rao-Wilton-Glisson (RWG) basis functions on triangular elements are employed to discretize the electric current [4]. In addition, the special basis and test functions derived from RWG basis function are introduced to ensure current continuity across the periodic boundaries [15]. The application of the Floquet-Bloch theorem reduces the computational domain of infinite periodic structures to a single unit cell but leads to the numerical evaluation of very slowly converging series.

2.2. Modified Ewald Transformation. We propose the following strategy to compute the spatial periodic Green’s function in an arbitrary point $\mathbf{u} = \mathbf{r} - \mathbf{r}'$.

If $u_z = |z - z'| > 0.5 \sqrt{A}$, PGF is computed by its spectral representation with exponential convergence, and thus no acceleration technique is required [16]. Consider

$$ G_p(\mathbf{r}, \mathbf{r}') = \sum_{m=-\infty}^{\infty} \sum_{n=-\infty}^{\infty} \frac{e^{j k_{mn} (\rho - \rho')}}{2 j A k_{mn}} e^{-j k_{mn} |z - z'|}, $$

(2)

where

$$ \mathbf{r} = \mathbf{r} + z \hat{z}, $$

$$ k_{mn} = k_{100} + \frac{2n}{A} \left[ m (\hat{a}_2 \times \hat{z}) + n (\hat{z} \times \hat{a}_1) \right]. $$

(3)

is the reciprocal lattice vector. Consider

$$ k_{mn} = \sqrt{k_0^2 - \mathbf{k}_{1mn} \cdot \mathbf{k}_{1mn}}, $$

(4)

where $\text{Re}(k_{mn}) \geq 0$, $\text{Im}(k_{mn}) < 0$.

If $u_z = |z - z'| < 0.5 \sqrt{A}$, PGF is computed by the representation of Ewald transformation [17], which are two exponential converging series. Consider

$$ G_p(\mathbf{r}, \mathbf{r}') = G_{p1}(\mathbf{r}, \mathbf{r}') + G_{p2}(\mathbf{r}, \mathbf{r}'), $$

(5)
\[
G_p(\vec{r}, \vec{r}') = \sum_{m=-\infty}^{\infty} \sum_{n=-\infty}^{\infty} e^{-j k_z m n} \frac{e^{-j k_m \cdot (\vec{r} - \vec{r}')}}{4 |\vec{k}_m^z|} + e^{j k_z m n} \frac{e^{j k_m \cdot (\vec{r} - \vec{r}')}}{4 |\vec{k}_m^z|},
\]

where \(G_p\) is the PGF, \(k_z\) is the wavenumber in the z-direction, and \(\vec{r}, \vec{r}'\) are the position vectors of the source and observation points, respectively.

2.3. Singularity Extraction. When the distance between the observation point \(\vec{r}\) and the source point \(\vec{r}'\) or one of its periodic extensions is electrically small, the contribution to Green’s function is essentially quasistatic. Usually the center distance between source and field triangles is less than 0.1\(\lambda\); then both are considered to be “near interaction.” Singular terms in (5) and (7) often arise, for example, from one of nine spatial terms \(R_{mn}\) with \(m, n = -1, 0\) or +1, depending on which term represents the source nearest to the observer \(\vec{r}\).

By extracting its singularity [18], the PGF can be expressed as
\[
G_{p1} = G_{p1}^0 + \frac{1}{4\pi R},
\]

where \(G_{p1}^0\) is the PGF of the primary source point and \(G_{p1}\) is the PGF of the source point nearest to the observer. The singularity subtraction technique [19] used in this paper can compute the singularity of \(G_p\).

Supposing the source point nearest to the observer point appears at \((1, 0)\) term rather than \((0, 0)\) term as shown in Figure 2, the primary source point can be transferred to \((1, 0)\) lattice. Then, the singular term always appears at \((1, 0)\) term.

\[
G_{p2}^0(\vec{r}, \vec{r}') = \frac{1}{4\pi R} + \frac{1}{4\pi R},
\]

where \(G_{p2}^0\) is the PGF of the secondary source point and \(G_{p2}\) is the PGF of the secondary source point nearest to the observer.
where $F^{00}_{p^2}$ is the nonsingular part

$$F^{00}_{p^2}(\vec{r}, \vec{r}') = \frac{1}{4\pi R} \cdot \text{real} \left\{ e^{-jk_0} \cdot \text{erfc} \left( RE - \frac{j k_0}{2E} \right) - \text{erfc} \left( -\frac{j k_0}{2E} \right) \right\}. \quad (10)$$

As $R \to 0$, the result of applying L'Hôpital's rule on (10) is as follows:

$$\lim_{R \to 0} F^{00}_{p^2}(\vec{r}, \vec{r}') = \frac{1}{2\pi} \cdot \text{real} \left[ -jk_0 \text{erfc} \left( -\frac{j k_0}{2E} \right) - \frac{2E}{\sqrt{\pi}} e^{k_0^2/4E^2} \right]. \quad (11)$$

2.4. Interpolation of PGF. PGFs are very smooth and amenable to interpolation. Fast interpolation of PGFs can greatly reduce the time of matrix filling. The reason is that interpolating the PGF values from the table is much faster than directly computing the Ewald sums (since this has to be done for every pair of source and field triangles in the numerical integration using cubature formulas). Trilinear interpolation [20] suited for the calculation of complex-valued function with very high efficiency is applied here. As shown in Figure 3, the cuboid grouping the unit cell is equally divided into many subgrids with side length $L$. Yellow points are the precomputed PGFs at the corner of the subgrid. Then the three-dimensional PGF values can be obtained by trilinear interpolation:

$$G(\vec{u}) = (1 - \alpha_x)(1 - \alpha_y)(1 - \alpha_z)G_{n_x+1,n_y+1,n_z+1} + (1 - \alpha_x)(1 - \alpha_y)\alpha_zG_{n_x+1,n_y+1,n_z+2} + (1 - \alpha_x)\alpha_y(1 - \alpha_z)G_{n_x+1,n_y+2,n_z+1} + \alpha_x(1 - \alpha_y)(1 - \alpha_z)G_{n_x+2,n_y+1,n_z+1} + \alpha_x(1 - \alpha_y)\alpha_zG_{n_x+2,n_y+1,n_z+2} + \alpha_x\alpha_y(1 - \alpha_z)G_{n_x+2,n_y+2,n_z+1} + \alpha_x\alpha_y\alpha_zG_{n_x+2,n_y+2,n_z+2},$$

where $\alpha_x = dx/L, \alpha_y = dy/L, \alpha_z = dz/L$.

The relative error $\varepsilon$ obtained by interpolating the PGFs is defined as

$$\varepsilon = \frac{|G^\text{int} - G^\text{ex}|}{G^\text{ex}}, \quad (13)$$

where $G^\text{int}$ and $G^\text{ex}$ are the interpolated and the exact data, respectively. The exact data are obtained by the above accelerating method with an accuracy $\varepsilon = 10^{-7}$. The relative errors are less than $\varepsilon = 10^{-7}$ and have, therefore, been neglected.

Take the multilayered structure with orthogonal lattices as an example. The relative error is shown in Figure 4 for the interpolation of PGF involving a series of interpolation points through the trilinear interpolation method, with various values of $N$ int. In Figure 4, positions $(A, \ldots, D)$ are the sampling points for interpolation. A very high level of accuracy is easily obtained with a relatively small number of interpolation points. The side length of the subgrid is usually fixed to 0.02$\lambda$, and the higher interpolation accuracy ($\varepsilon < 10^{-7}$) can be obtained.

3. ACA Technique

Let the $m \times n$ rectangular matrix $Z_{mn \times n}$ represent the coupling between two well-separated groups in the MoM computation. The ACA algorithm aims to accurately approximate $Z_{mn \times n}$ by a low-rank matrix $\tilde{Z}_{mn \times n}$. In particular, the ACA algorithm
constructs the approximate matrix $\bar{Z}_{m\times n}$ through a product form. Namely,

$$\bar{Z}_{m\times n} = U_{m\times r} V_{r\times n} = \sum_{i=1}^{r} u_{m\times i}^i v_{i\times n}^i,$$  \hspace{1cm} (14)

where $r$ is the rank of $\bar{Z}_{m\times n}$ or the effective rank of the matrix $Z_{m\times n}$; $U_{m\times r}$ and $V_{r\times n}$ are two dense rectangular matrices; $u_{m\times i}^i$ is the column $i$th of $U$; and $v_{i\times n}^i$ is the row $i$th of $V$, respectively. The goal of ACA is to achieve

$$\| Z_{m\times n} - \bar{Z}_{m\times n} \|_F \leq \epsilon \| Z_{m\times n} \|_F$$     \hspace{1cm} (15)

for a given tolerance $\epsilon$. Note that in this paper $\| \cdot \|$ refers to the matrix Frobenius norm $[9]$.

The ACA technique is numerically based on the principle of the approximation above but does not need to know all terms of $Z_{m\times n}$. Indeed, only $r$ rows and $r$ columns are calculated during the algorithm allowing a significant reduction in the number of integral computations to perform. Therefore, not only is the memory requirement reduced but the assembly time is also saved. Yet, if we agree to deal with the MVP $U_{m\times n} V_{r\times n} h_{i\times 1}$ instead of the exact value of $Z_{m\times n} h_{i\times 1}$, the number of operations needed is also remarkably reduced by performing $U_{m\times r} V_{r\times n} h_{i\times 1}$. Thus, if an iterative solver is used, one can accelerate matrix vector multiply and save once again the computation time $[10]$.

The compression rate $\eta$ of matrix block $Z_{m\times n}$, which denotes the memory saved by using ACA, can be expressed as

$$\eta \% = \left(1 - \frac{r \cdot (m + n)}{m \cdot n}\right) \cdot 100\%.$$ \hspace{1cm} (16)

Obviously, the approximation by ACA approach above is useful only if $r < (1/2) \cdot \min[m, n]$, where $\bar{Z}_{m\times n}$ presents similar values. For the electromagnetic application, the effective rank $r < \min(m, n)$. Therefore, instead of storing entire $m \times n$ entries, the algorithm only requires storing $(m + n) \times r$ entries.

4. Numerical Results

The EFIE and ACA approaches to PEC are implemented using Fortran-95 language. Numerical experiments run on a PC with Intel i5-2400 3.1 GHz CPU and 16 GB RAM. The resulting impedance matrices are iteratively solved by using the transpose-free quasi-minimal residual (TFQMR) solver with diagonal preconditioning $[21]$; the relative error tolerance is set to be $10^{-3}$.

4.1. Infinite Metallic Rod. As a 2D example, we study the scattering properties of a photonic band gap (PBG) structure consisting of infinite metallic rod $[17]$. Each unit cell consists of two infinitely long metallic rods. The inset of Figure 5 is two meshed cylinders in a unit cell. The axis of each cylinder is along $x$-direction: $\vec{a}_1 = \vec{x}$ 6 mm and $\vec{a}_2 = \vec{y}$ 6 mm. Each cylinder has the radius 0.6 mm and length 6 mm. The space between two cylinders is 6 mm. Each cylinder was discretized into 494 triangle facets with 741 unknowns to ensure accurate results obtained in the entire frequency band. Figure 5 shows the magnitude of the transmission and reflection coefficients for the Floquet TM$_{00}$ mode with the electric field oriented in the $x$-direction. Our results agree well with the reference results obtained using 2D IE approach $[15]$. The results converged for 9 terms in both Ewald sums and there are 1482 unknowns for the geometry. The time for matrix filling and solving is 0.58 s and 0.21 s per frequency point, respectively. In ACA, the box size is $6 \times 6 \times 6$ mm$^3$; that is, each box includes

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure4.png}
\caption{The relative error of a series of points. Reference structure: primitive vector $|\vec{a}_1| = |\vec{a}_2| = 0.55\lambda$, $h = 2.2\lambda$; phase shifts $k_x = k \cos(45^\circ), k_y = k \sin(30^\circ)$. Accuracy reached interpolating PGFs through trilinear interpolation, with a 3D grid of $N_{\text{int}}^x \times N_{\text{int}}^y \times 2N_{\text{int}}^z$ equispaced points in $-0.55\lambda < \Delta x < 0.55\lambda$, $-0.55\lambda < \Delta y < 0.55\lambda$, $0 < \Delta z < 2.2\lambda$. The data plotted refers to the straight line from $(0.15\lambda, 0.15\lambda, 0)$ to $(0.15\lambda, 0.15\lambda, 2.2\lambda)$. The number of equally spaced sampling points is 4.}
\end{figure}
one cylinder. If the trilinear interpolation method is not used, the time of matrix filling would increase dramatically to 128.6 s. Therefore, trilinear interpolation of PGFs can reduce the time of matrix filling by more than 99.5%.

Next, the number of layers increases from 2 to 5, 10 and 21, respectively. Figure 6 shows the transmission coefficient of multilayered structure. The transmission coefficient curves exhibit the typical resonances of photonic band gap materials. It can be seen that the null of transmission becomes deeper as the layer number increases. The total solution times for per frequency point are 5.1 s, 12.4 s, and 32.5 s, respectively.

The 21-layer structure is meshed into 10374 triangle facets with 15561 unknowns. The corresponding times of direct solution by MoM without ACA acceleration are 187.6 s and 75.2 s per frequency point, respectively. Obviously, ACA can greatly reduce the total solution time, especially for frequency sweeping.

### 4.2. Woodpile
A 3D example showing the inset in Figure 7 is a woodpile MPC with $w = 200\,\text{nm}$, $h = 300\,\text{nm}$ and in-layer lattice constant $d = 1\,\mu\text{m}$. The incident wave is a TE polarized (i.e., $E$ parallel to the top metallic layer) plane wave through the stacking direction. The number of layers are 4, 8, and 16, respectively. Two intersecting metallic pipes are considered to be fully contacting with each other in numerical modeling. The ACA box size is $1 \times 1 \times 0.3\,\mu\text{m}^3$. The reflection coefficients are shown in Figure 7. Almost the same result is obtained for TM polarization. Our numerical results agree well with that of Ansys HFSS. Previous results reported by other studies [12, 22] for a similar kind of the structural geometry also validate our calculated results. For the 16-layer MPC with 13149 unknowns (8766 triangle facets), the times for matrix filling and linear system solving are 40.3 s and 15.2 s per frequency point, respectively. The corresponding times of direct solution by MoM without ACA acceleration are 132.3 s and 92.5 s, respectively.

### 5. Conclusion
An integral equation accelerated with adaptive cross approximation (ACA) is proposed for the analysis of multilayered metallic photonic crystal. The ACA is a purely algebraic and
kernel independent algorithm. It is verified to be suited for accelerating integral equation analysis of periodic structure. The ACA can be applied to compress impedance matrix blocks and accelerate MVP for reducing computational complexity. A modified Ewald transformation and an efficient interpolation method are also proposed to fast compute the periodic Green’s function; hence, that can greatly reduce the time of matrix filling. Numerical results confirm the validity and accuracy of the proposed method. In our next stage of research, ACA will be applied to accelerate hybrid field integral equation for arbitrarily complex multilayered periodic structure composed of metal and dielectric materials with more unknowns.

**Conflict of Interests**

The authors declare that there is no conflict of interests regarding the publication of this paper.

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