Discrete-dipole approximation for scattering calculations

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The discrete-dipole approximation (DDA) for scattering calculations, including the relationship between the DDA and other methods, is reviewed. Computational considerations, i.e., the use of complex-conjugate gradient algorithms and fast-Fourier-transform methods, are discussed. We test the accuracy of the DDA by using the DDA to compute scattering and absorption by isolated, homogeneous spheres as well as by targets consisting of two contiguous spheres. It is shown that, for dielectric materials (|\varepsilon| \leq 2), the DDA permits calculations of scattering and absorption that are accurate to within a few percent.

1. INTRODUCTION

The discrete-dipole approximation (DDA) is a flexible and powerful technique for computing scattering and absorption by targets of arbitrary geometry. The development of efficient algorithms and the availability of inexpensive computing power together have made the DDA the method of choice for many scattering problems. Bohren and Singham\(^1\) reviewed recent advances in calculations of backscattering by nonspherical particles, including DDA results. In this paper we review the DDA, with particular attention to recent developments.

In Section 2 we briefly summarize the conceptual basis for the DDA. The relation of the DDA to other finite-element methods is discussed.

DDA calculations require choices for the locations and the polarizabilities of the point dipoles that are to represent the target. In Section 3 we discuss these choices, with attention to the important question of dipole polarizabilities. Criteria for the validity of the DDA are also considered.

Recent advances in numerical methods now permit the solution of problems involving large values of \(N\), the number of point dipoles. These developments, particularly the use of complex-conjugate gradient (CCG) methods and fast-Fourier-transform (FFT) techniques, are reviewed in Section 4.

We illustrate the accuracy of the method in Section 5 by using the DDA to compute scattering by a single sphere and by targets consisting of two spheres in contact and by comparing our results with the exact results for these geometries. Although these are only examples, they make clear that the DDA can be used to compute highly accurate results for targets with dielectric constants that are not too large (|\varepsilon| \leq 2).

A portable FORTRAN implementation of the DDA, the program DDSCAT, has been developed by the authors and is freely available.\(^2\)

2. WHAT IS THE DISCRETE-DIPOLE APPROXIMATION?

A. Approximation

Given a target of arbitrary geometry, we seek to calculate its scattering and absorption properties. Exact solutions to Maxwell's equations are known only for special geometries such as spheres, spheroids, or infinite cylinders, so approximate methods are in general required. The DDA is one such method.

The basic idea of the DDA was introduced in 1964 by DeVoe,\(^3\) who applied it to study the optical properties of molecular aggregates; retardation effects were not included, so DeVoe's treatment was limited to aggregates that were small compared with the wavelength. The DDA, including retardation effects, was proposed in 1973 by Purcell and Pennypacker,\(^5\) who used it to study interstellar dust grains. Simply stated, the DDA is an approximation of the continuum target by a finite array of polarizable points. The points acquire dipole moments in response to the local electric field. The dipoles of course interact with one another via their electric fields,\(^6\) so the DDA is also sometimes referred to as the coupled dipole approximation.\(^7\) The theoretical basis for the DDA, including radiative reaction corrections, is summarized by Draine.\(^6\)

Nature provides the physical inspiration for the DDA: in 1909 Lorentz showed\(^9\) that the dielectric properties of a substance could be directly related to the polarizabilities of the individual atoms of which it was composed, with a particularly simple and exact relationship, the Clausius–Mossotti (or Lorentz–Lorenz) relation, when the atoms are located on a cubic lattice. We may expect that, just as a continuum representation of a solid is appropriate on length scales that are large compared with the interatomic spacing, an array of polarizable points can accurately approximate the response of a continuum.
target on length scales that are large compared with the interdipole separation.

For a finite array of point dipoles the scattering problem may be solved exactly, so the only approximation that is present in the DDA is the replacement of the continuum target by an array of $N$-point dipoles. The replacement requires specification of both the geometry (location $r_j$ of the dipoles $j = 1, \ldots, N$) and the dipole polarizabilities $\alpha_j$.

For monochromatic incident waves the self-consistent solution for the oscillating dipole moments $\mathbf{p}_j$ may be found, as is discussed in Section 4 below; from these $\mathbf{p}_j$ the absorption and scattering cross sections are computed. If DDA solutions are obtained for two independent polarizations of the incident wave, then the complete amplitude-scattering matrix can be determined.

With the recognition that the polarizabilities $\alpha_j$ may be tensors, the DDA can readily be applied to anisotropic materials. The extension of the DDA to treat materials with nonzero magnetic susceptibility is also straightforward, although for most applications magnetic effects are negligible.

**B. Relation to Other Methods**

The DDA is one particular discretization method for solving Maxwell’s equations in the presence of a target. The DDA was anticipated in the engineering literature by the method of moments, which is a general prescription for discretization of the integral form of Maxwell’s equations. There is a close correspondence between the DDA and discretizations that are based on the digitized Green’s function (DGF) method or the volume-integral-equation formulation (VIEF). In the low-frequency limit $kd \to 0$ (where $d$ is the interdipole spacing or the side of a cubic subvolume and $k = \omega/c$), it has been shown that the DDA is indeed equivalent to the DGF/VIEF. However, researchers developing the DGF/VIEF approach encountered difficulties in trying to calculate $O((kd)^2)$ corrections, whereas the DDA, because the fundamental approximation is simple and physically clear, permits rigorous derivation of the $O((kd)^2)$ correction (see Subsection 3.B below). On the other hand, the integral form of Maxwell’s equations often serves as a convenient point of departure, and there is now a strong interplay between both approaches.

The principal limitation of the DDA involves the handling of target boundaries, since the geometry of the DDA array has a minimum length scale that is equal to the interdipole spacing $d$. For targets with large values of the refractive index $m$, the accuracy of the DDA suffers. In principle we can make the DDA as accurate as we desire by increasing $N$, the number of dipoles representing the target (thus decreasing $d$), but very large values of $N$ become computationally prohibitive.

**3. SPECIFICATION OF THE DIPOLE ARRAY**

**A. Geometry**

Each dipole may be thought of as representing the polarizability of a particular subvolume of target material. If we wished to approximate a certain target geometry (e.g., a sphere) with a finite number of dipoles, then we might consider using some number of closely spaced, weaker dipoles in regions near the target boundaries to do a better job of approximating the boundary geometry. Some researchers have followed this approach, although it clearly offers benefits, there is also a substantial cost: as we show below, FFT methods can greatly accelerate the computations that are required for solving the scattering problem, but only if the dipoles are located on a periodic lattice. With FFT methods we can perform calculations (on a workstation) for numbers $N$ of dipoles (e.g., $10^5$) that are much greater than the largest values ($\sim 10^4$) that could readily be handled without FFT methods. We therefore elect to use FFT methods, accepting the requirement that the dipoles be located on a periodic lattice. We further restrict ourselves to cubic lattices.

This option still permits considerable latitude in representation of the continuum target. For a given target geometry we use the following algorithm to generate the dipole array. We assume that the target orientation is fixed relative to a coordinate system $\hat{x}, \hat{y}, \hat{z}$.

1. Generate a trial lattice that is defined by the lattice spacing $d$ and coordinates $(x_0, y_0, z_0)$ of the lattice point nearest the origin.

2. Let the array now be defined as all lattice sites located within the volume $V$ of the continuum target.

3. Try different values of $d$ and $(x_0, y_0, z_0)$ and maximize some goodness-of-fit criterion, subject to the constraint that the number of points in the dipole array not be so large as to be computationally prohibitive. The goodness-of-fit criteria are basically geometric in nature, as well as somewhat arbitrary, and are not discussed here.

4. We now have a list of occupied sites $j = 1, \ldots, N$. We imagine that each of these occupied sites represents a cubic subvolume $d^3$ of material centered on the site. In the current implementation of DDSCAT we do not distinguish between lattice sites near the surface and those in the interior. We therefore rescale the array by requiring that $d = (V/N)^{1/3}$, so that the volume $Nd^3$ of the occupied lattice sites is equal to the volume $V$ of the original target.

5. For each occupied site $j$ assign a dipole polarizability $\alpha_j$.

**B. Dipole Polarizabilities**

There has been some controversy concerning the best method of assigning the dipole polarizabilities. Purcell and Pennypacker used the Clausius–Mossotti polarizabilities:

$$\alpha_j^{CM} = \frac{3d^3}{4\pi} \frac{\varepsilon_j - 1}{\varepsilon_j + 2}$$

where $\varepsilon_j$ is the dielectric function of the target material at location $r_j$. As is well known, for an infinite cubic lattice the Clausius–Mossotti prescription is exact in the dc limit $kd \to 0$. Draine and Goedecke and O’Brien showed that the polarizabilities should also include a radiative reaction correction of $O((kd)^2)$; the same correction term was independently proposed by Iskander et al. and by Hage and Greenberg. Unfortunately the derivation of this correction assumed that the electric
field could be taken to be uniform over cubical regions
of volume \( d^3 \), and this assumption itself introduces errors
of \( O([kd]^{3/2}) \). Another approach was taken by Dungey
and Bohren,\(^{27}\) who arrived at a different \( O([kd]^{3/2}) \)
correction by requiring that each point dipole have the same
polarizability as a finite sphere of diameter \( d \), but with a
modified dielectric constant.

To clarify this issue Draine and Goodman\(^{26}\) considered
the following problem: for what polarizability \( \alpha(\omega) \)
will an infinite lattice of polarizable points have the
same dispersion relation as a continuum of refractive
index \( m(\omega) \)? The lattice dispersion relation (LDR)
may be found analytically; in the long-wavelength limit
\( kd \ll 1 \), the polarizability \( \alpha(\omega) \) is given as a series
expansion in powers of \( kd \) and \( m^2 = \varepsilon \):

\[
\alpha_{\text{LDR}} = \frac{\alpha_{\text{CM}}}{1 + (\alpha_{\text{CM}}/d^3)[(b_1 + m^2 b_2 + m^2 b_3 S)(kd)^2 - (2/3)i(kd)^3]},
\]

\[
b_1 = -1.891531, \quad b_2 = 0.1648469, \quad b_3 = -1.7700004, \quad S = \sum_{j=1}^{3} (\delta_i j \varepsilon_j)^2,
\]

where \( \alpha \) and \( \varepsilon \) are unit vectors defining the incident
direction and the polarization state. The \( O([kd]^{3/2}) \) term
in the LDR expansion differs from previously proposed corrections.\(^{18,20,21,27}\) The LDR prescription for \( \alpha(\omega) \) is,
by construction, optimal for wave propagation on an infinite
lattice, and it is reasonable to assume that it will
also be a good choice for finite dipole arrays. Extensive
DDA calculations for spheres, comparing different
prescriptions for the dipole polarizability, confirm that
the LDR prescription appears to be best for \( |m|kd < 1 \).\(^{26}\)
The different prescriptions have also been compared in
calculations of scattering by two touching spheres\(^{26}\) (see
Fig. 9 below).

C. Validity Criteria

There are two obvious criteria for validity of the DDA:
(1) \( |m|kd \ll 1 \) (so that the lattice spacing \( d \) is small compared
with the wavelength of a plane wave in the target
material), and (2) \( d \) must be small enough (\( N \) must be
large enough) to describe the target shape satisfactorily.

Define the effective radius \( a_{\text{eff}} \) of a target of volume \( V \)
by \( a_{\text{eff}} = (3V/4\pi)^{1/3} \). The first criterion is then equivalent to

\[
N > (4\pi/3)|m|^3(a_{\text{eff}})^3.
\]

Thus targets with large values of \( |m| \) or scattering
problems with large values of \( a_{\text{eff}} \) will require that large
numbers of dipoles be used to represent the targets.

Unfortunately the second criterion has not yet been
formulated precisely. Draine\(^8\) shows that even in the
\( kd \to 0 \) limit the polarizations are too large in the surface
monolayer of dipoles in a pseudosphere, and similar errors
must also occur for other target shapes. As a result the
rate of energy absorption by the dipoles in the surface
monolayer is too large, which leads to an error in the
overall absorption cross section in proportion to the fraction
\( \sim N^{-1/3} \) of the total volume that is contributed by the

surface monolayer. As a result, when \( |m| > 1 \) the DDA
can seriously overestimate absorption cross sections, even
in the dc limit \( |m|kd \ll 1 \). For materials with \( |m| \gg 1 \) it
appears that other techniques (e.g., the method of Rouleau
and Martin\(^{26}\)) may be superior to the DDA.

4. COMPUTATIONAL CONSIDERATIONS

A. Scattering Problem

The electromagnetic scattering problem must be solved
for the target array of point dipoles \( (j = 1, \ldots, N) \) with
polarizabilities \( a_j \), located at positions \( \mathbf{r}_j \). Each dipole
has a polarization \( \mathbf{P}_j = a_j \mathbf{E}_j \), where \( \mathbf{E}_j \) is the electric
field at \( \mathbf{r}_j \) that is due to the incident wave \( \mathbf{E}_{\text{inc},j} =
\mathbf{E}_0 \exp[i(k \mathbf{r}_j - io t)] \) plus the contribution of each of the

other \( N - 1 \) dipoles:

\[
\mathbf{E}_j = \mathbf{E}_{\text{inc},j} - \sum_{\mathbf{k} \neq j} A_{jk} \mathbf{P}_k,
\]

where \( -A_{jk} \mathbf{P}_k \) is the electric field at \( \mathbf{r}_j \) that is due to
dipole \( \mathbf{P}_k \) at location \( \mathbf{r}_k \), including retardation effects.
Each element \( A_{jk} \) is a \( 3 \times 3 \) matrix:

\[
A_{jk} = \frac{\exp(ikr_{jk})}{r_{jk}} \times \left[ \begin{array}{c}
\kappa^2 (\mathbf{r}_{jk} \cdot \mathbf{\hat{r}}_{jk} - 1_3) + \frac{ikr_{jk} - 1}{r_{jk}^2} (3\mathbf{r}_{jk} \cdot \mathbf{\hat{r}}_{jk} - 1_3) \end{array} \right],
\]

\[
j \neq k,
\]

where \( \kappa = \omega/c, r_{jk} = |\mathbf{r}_j - \mathbf{r}_k|, \mathbf{\hat{r}}_{jk} = (\mathbf{r}_j - \mathbf{r}_k)/r_{jk} \),
and \( 1_3 \) is the \( 3 \times 3 \) identity matrix. Defining \( A_{jj} = \alpha_j^{-1} \)
reduces the scattering problem to finding the polarizations
\( \mathbf{P}_j \) that satisfy a system of \( 3N \) complex linear equations:

\[
\sum_{k=1}^{N} A_{jk} \mathbf{P}_k = \mathbf{E}_{\text{inc},j}.
\]

Once Eq. (7) has been solved for the unknown polarizations
\( \mathbf{P}_j \), the extinction and absorption cross sections \( C_{\text{ext}} \)
and \( C_{\text{abs}} \) may be evaluated:\(^6\)

\[
C_{\text{ext}} = \frac{4\pi k}{|\mathbf{E}_0|^2} \sum_{j=1}^{N} \text{Im}(\mathbf{E}_{\text{inc},j}^* \cdot \mathbf{P}_j),
\]

\[
C_{\text{abs}} = \frac{4\pi k}{|\mathbf{E}_0|^2} \sum_{j=1}^{N} \left[ \text{Im}(\mathbf{P}_j \cdot (\alpha_j^{-1})^* \mathbf{P}_j^*) - \frac{2}{3} k^3 |\mathbf{P}_j|^2 \right].
\]

The scattering cross section \( C_{\text{scat}} = C_{\text{ext}} - C_{\text{abs}} \). Differential scattering cross sections may also be directly
evaluated once the \( \mathbf{P}_j \) are known.\(^8\) In the far field the
scattered electric field is given by

$$E_{\text{scat}} = \frac{\hbar^2}{r} \exp(-ikr) \sum_{j=1}^{N} \exp(-ik\vec{r} \cdot \vec{r}_j)(\vec{r} - \vec{r}_j)P_j.$$  \hspace{1cm} (10)

The problem is that matrix $A$ is large and full, i.e., a matrix that in general has few zero elements. We would like (1) to solve the linear system of Eq. (7) efficiently and (2) to solve efficiently for multiple cases of the incident plane wave $E_{\text{inc}}$. LU decomposition\(^{39}\) is the method of choice for small problems. It solves problem (2) because once the LU decomposition has been obtained the solution for a new right-hand side $E_{\text{inc}}$ requires just one matrix–vector multiplication. However, it is not feasible or efficient for large problems because of the need to store $A$ and the LU decomposition and because of computer time that is proportional to $N^3$.

For homogeneous, rectangular targets the matrix $A$ has block-Toeplitz symmetry,\(^{31}\) and algorithms exist for finding $A^{-1}$ in $O(N^2)$ operations rather than in the $O(N^3)$ operations that are required for general matrix inversion. It seems possible that the fundamental symmetries of $A$ may permit efficient algorithms for finding $A^{-1}$ even for targets that are inhomogeneous or nonrectangular.

### B. Complex-Conjugate Gradient Method

Rather than direct methods for solving Eq. (7), CCG methods for finding $P$ iteratively have proven effective and efficient. The particular CCG algorithm that is used in DDISCATT is described by Draine.\(^{6}\) As $P$ has $3N$ unknown elements, CCG methods in general are only guaranteed to converge in $3N$ iterations. In fact, however, $\sim 10-10^2$ iterations are often found to be sufficient to obtain a solution to high accuracy.\(^{6,32}\) The choice of conjugate-gradient variant may influence the convergence rate.\(^{33-35}\)

When $N$ is large, CCG methods are much faster than are direct methods for finding $P$. The fact that CCG methods in practice converge relatively rapidly is presumably a consequence of the basic symmetries of $A$.

As an iterative technique the CCG requires an initial guess for $P$. The simplest choice is $P = 0$. Attempts to improve on this by use of the scattering-order approximation\(^{36}\) as a first estimate for $P$ were found to offer little, if any, advantage over starting with $P = 0$.\(^6\)

The iterative method that is described here must be repeated for each incident plane wave (cf. Eq. (7)). Notice that if we had $A^{-1}$ then for each new incident wave $E_{\text{inc}}$ the solution $P = A^{-1}E_{\text{inc}}$ would be obtained from a single matrix–vector multiplication. In principle we can recover eigenvalues and eigenvectors from conjugate-gradient iteration.\(^{37,38}\) Some attempts to improve the efficiency of the conjugate-gradient method for multiple incident electric fields have been reported.\(^{39-41}\) FORTAN implementations of approximately 15 CCG methods are available from us.\(^{42}\)

### C. Fast Fourier Transforms

The computational burden in the CCG method primarily consists of matrix–vector multiplications of the form $A \cdot v$. Goodman et al.\(^{32}\) show that the structure of the matrix $A$ implies that such multiplications are essentially convolutions, so that FFT methods can be employed to evaluate $A \cdot v$ in $O(N \ln N)$ operations rather than in the $O(N^2)$ operations that are required for general matrix–vector multiplication. Since $N$ is large this is an important calculational breakthrough. As mentioned above, FFT methods require that the dipoles be situated on a periodic lattice, which is most simply taken to be cubic.

FFT methods in effect require the computation of three-dimensional FFT’s over $8N_xN_yN_z$ points, where $N_xN_yN_z$ is the number of sites in a rectangular region of the lattice containing all the $N$ occupied lattice sites. Therefore, in the case of a fractal structure with a large volume-filling factor for vacuum, FFT methods may lose some of their advantage over conventional techniques for evaluating $A \cdot v$, since $N_xN_yN_z$ may be much larger than the actual number of dipoles $N$.

We note that CCG and FFT techniques are routinely used for numerical solutions of electromagnetic problems in engineering.\(^{34,43,44}\)

### D. Memory and CPU Requirements

If the $N$ dipoles were located at arbitrary positions $r_j$, then the $9N^2$ elements of $A$ would be nondegenerate. Storage of these elements (with 8 bytes/complex number) would require $72(N/10^3)^2$ Mbytes. By locating the dipoles on a lattice, the elements of $A$ become highly degenerate, since they depend only on the displacement $r_j - r_i$. As a result the memory requirements depend approximately linearly on $N$ rather than on $N^2$. The program DDISCATT has a total memory requirement of $\sim 0.58(N_xN_yN_z/1000)$ Mbytes, where $N_x \times N_y \times N_z$ is the rectangular volume containing all the $N$ dipoles. Thus for a target fitting into a $32 \times 32 \times 32$ portion of the lattice (e.g., an $N = 17,904$ pseudosphere), DDISCATT requires $\sim 19$ Mbytes of memory. We therefore see that memory requirements begin to be a consideration for targets that are much larger than $10^4$ dipoles.

The CPU requirements also are significant for large targets. On a Sun 4/50 (Sparcstation IPX), a single CCG iteration requires $\sim 3.0(N_xN_yN_z/10^3)$ CPU s; thus one iteration for a $N = 17,904$ pseudosphere requires $\sim 100$ CPU s. Between 10 and $10^3$ iterations are typically necessary to solve for a single incident direction and polarization; thus orientational averaging with the DDA can be time consuming if many target orientations are required. It is for this reason that $T$-matrix methods,\(^{45,46}\) which exploit efficient procedures for orientational averaging,\(^{47}\) are competitive for some target geometries as well as recursive $T$-matrix algorithms currently being developed by Chew and Lu\(^{48}\) and Chew et al.\(^{42}\)

### 5. ACCURACY OF THE DISCRETE-DIPOLE APPROXIMATION

At this time there is no known way to predict precisely the accuracy of DDA calculations. Instead we rely on examples to guide us. Spheres are most convenient since exact solutions are readily available for comparison. Note that the DDA does not give preferential treatment to any geometry (with the possible exception of rectangular targets), so spheres constitute a representative test of
the accuracy of the DDA. Clusters of spheres, for which exact solutions can now also be computed, are also useful.

Several earlier studies have examined the accuracy of the DDA,6,8,9,37,50,51 but only one6 employed the LDR polarizabilities. The DDA is most accurate for targets with refractive indices \( m \) near unity, with the accuracy declining (for fixed \( ka_{\text{eff}} \) and \( N \)) as \( |n - 1| \) increases. Here we employ the LDR polarizabilities and concentrate on two cases: \( m = 1.33 + 0.01i \), representative of moderately polarizable material, and \( m = 2 + i \), representative of a strongly polarizable, strongly absorbing material in the visible or the ultraviolet. As is well known, very large refractive indices \( (|m| > 1) \) occur for conducting materials in the infrared; the DDA is not particularly well suited for this regime, and we do not consider any such examples here. We will examine the dependence of the accuracy on both \( ka_{\text{eff}} \) and \( N \).

A. Spheres

1. Total Cross Sections

For a given incident wave the scattering problem is solved by CCG iteration until a specified level of accuracy is achieved. All the results shown here are converged until \( |\mathbf{A} \cdot \mathbf{P} - E_{\text{inc}}| < 10^{-6} |E_{\text{inc}}| \). Figures 1 and 2 show the errors in the absorption and scattering efficiency factors

\[
Q_{\text{abs}} = C_{\text{abs}}/\pi a^2 \quad \text{and} \quad Q_{\text{sca}} = C_{\text{sca}}/\pi a^2, \quad \text{as functions of} \quad x = 2\pi a/\lambda, \quad \text{for pseudospheres containing} \quad N \quad \text{dipoles; the curves are labeled by} \quad N. \quad \text{Each curve terminates at the point at which} \quad |m|kd = 1. \quad \text{The pseudospheres are obviously not precisely spherically symmetric; the DDA calculations in Figs. 1–8 are for radiation incident along the (1,1,1) direction. Figure 1 shows results for} \quad m = 1.33 + 0.01i \quad (\varepsilon = 1.769 + 0.0266i). \quad \text{It is clear that for a modest refractive index, such as} \quad m = 1.33 + 0.01i, \quad Q_{\text{sca}} \quad \text{and} \quad Q_{\text{abs}} \quad \text{may be obtained to} \quad \approx 2\% \quad \text{accuracy provided that} \quad N \quad \text{is large enough that the criterion} \quad |m|kd < 1 \quad \text{is satisfied.}

Figure 2 shows results for \( m = 2 + i \quad (\varepsilon = 3 + 4i) \). In this case the larger refractive index leads to larger errors, but the errors in \( Q_{\text{sca}} \) are still quite small. Somewhat surprisingly the largest errors tend to occur in the Rayleigh limit \( ka \ll 0.5 \); here \( Q_{\text{sca}} \) is in error by \( \approx 3\% \), and \( Q_{\text{abs}} \) is in error by \( 5\% \) for \( N = 1064 \). For fixed \( ka \) the errors decline as \( N \) is increased. It is apparent that even for a refractive index \( m = 2 + i \), which is moderately large and highly absorptive, scattering and absorption cross sections can be calculated to accuracies of \( 1\% \) and \( 4\% \), respectively, for pseudospheres containing \( \approx 10^4 \) dipoles.

2. Differential Scattering

Figures 1 and 2 show that the total scattering cross section can be calculated fairly accurately. Differential scattering efficiencies \( dQ/d\Omega = S_{11}/\pi (ka)^2 \) for unpolarized light incident upon a sphere are shown in Figs. 3–8 for two refractive indices \( (m = 1.33 + 0.01i \text{ and} \quad m = 2 + i) \) and \( ka = 3, 5, \text{ and} 7 \). The DDA results are shown for different numbers of dipoles \( N \). The lower panel in Figs. 3–8 displays the fractional error:

\[
\text{error} = \frac{dQ/d\Omega(\text{DDA}) - dQ/d\Omega(\text{Mie})}{dQ/d\Omega(\text{Mie})}. \quad (11)
\]

The fractional error becomes quite large for scattering angle \( \theta > 150^0 \) because \( dQ/d\Omega \) itself becomes quite small for backscattering; hence small absolute errors in \( dQ/d\Omega \) correspond to large fractional errors.

From Figs. 1–8 we may draw the following conclusions:

1. For pseudospheres with \( |m| \leq 3 \), total scattering and absorption can be computed to accuracies of a few
B. Two Contiguous Spheres

Recent investigations have studied the general solution to Maxwell's equations for clusters of spheres.\textsuperscript{52,53} The availability of this exact solution offers a unique opportunity to test the accuracy of the DDA when it is applied to nonspherical targets.\textsuperscript{53,54}

Figure 9 presents a comparison of a two-sphere, multipole solution with the DDA as a function of $x = ka_{\text{eff}}$. Results are shown for three different prescriptions for the dipole polarizabilities: CMRR (Clausius–Mossotti plus radiative reaction), DGF–VIEF, and LDR. The refractive index is $m = 1.33 + 0.01i$, and the dipoles are placed on a $64 \times 32 \times 32$ cubic lattice. The calculations are for percent if the validity criterion [relation (4)] is satisfied, and $N \approx 10^3$, so that the surface geometry is reasonably well approximated.

2. For pseudospheres, if the validity criterion [relation (4)] is satisfied, differential scattering cross sections can be calculated to reasonable accuracy.

For $m = 1.33 + 0.01i$, fractional errors in $dQ_{\text{scat}}/d\Omega$ are within $\sim 20\%$ for all scattering directions, with rms errors of $\leq 6\%$. For $m = 2 + i$, fractional errors in $dQ_{\text{scat}}/d\Omega$ are within $\sim 30\%$ for all scattering directions, with rms values of $\leq 11\%$. 

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Fig. 3. Differential scattering cross section for pseudospheres with $m = 1.33 + 0.01i$ and $ka = 3$. The curves are labeled by $N$, the number of dipoles in the pseudosphere.

Fig. 4. Same as Fig. 3 but for $ka = 5$.

Fig. 5. Same as Fig. 3 but for $ka = 7$.

Fig. 6. Same as Fig. 3 but for $m = 2 + i$. 

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References:

\textsuperscript{52,53} Maxwell's equations for clusters of spheres.\textsuperscript{52,53}

\textsuperscript{53,54} The availability of this exact solution offers a unique opportunity to test the accuracy of the DDA when it is applied to nonspherical targets.\textsuperscript{53,54}

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6. RECENT RESEARCH

The DDA has recently been applied to a number of diverse problems, including (since 1990) calculations of scattering of unpolarized light incident at an angle $\alpha = 30^\circ$ relative to the axis passing through the centers of the two spheres. As is noted in Section 3, the LDR method for specifying the polarizabilities results in very accurate values for $Q_{\text{sc}}$ and $Q_{\text{abs}}$ and is clearly preferable to the CMRR and DGF/VIEF prescriptions.

In Fig. 10 we compare the exact solution for scattering by two touching spheres (solid curve) with the solution derived from the DDA ($\times$’s) for $m = 1.33 + 0.001i$ and $ka_{\text{eff}} = 10$. The exact results for $S_{11}$ and $S_{22}$ are compared with results that were computed with the DDA. The overall agreement is very good, although errors can be seen for scattering angle $\theta \approx 50^\circ$.
and absorption by rough and porous particles, interstellar graphite particles, aggregate particles, inhomogeneous particles, structures on surfaces, microwave scattering by ice crystals, and single scattering properties of ice crystals.

7. SUMMARY

The discrete-dipole approximation, or the coupled-dipole method, is a flexible and powerful tool for computing scattering and absorption by arbitrary targets. For targets with |m| ≤ 2 (with m being the complex refractive index), scattering and absorption cross sections can be evaluated to accuracies of a few percent provided that the number N of dipoles satisfies relation (4). The use of CGC and FFT algorithms permits calculations for N as large as ~10^5, so that scattering problems with kα eff ≈ 10 can be studied with scientific workstations.

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