Magnetic Nanoantennas Made of Plasmonic Nanoclusters for Photoinduced Magnetic Field Enhancement

Mahsa Darvishzadeh-Varcheie, Caner Guclu, and Filippo Capolino

Department of Electrical Engineering and Computer Science, University of California, Irvine, California 92697, USA

(Received 8 March 2017; revised manuscript received 5 July 2017; published 31 August 2017)

We focus on a category of nanoantennas called magnetic nanoantennas made of a circular cluster of gold nanospheres that leads to enhanced local magnetic field oscillating at optical frequency. We elaborate on the magnetic field enhancement and the magnetic-to-electric-field ratio, i.e., the local field admittance, when the nanoantenna is illuminated by a single plane wave and by superposition of two plane waves to maximize the magnetic-only response. Single-dipole approximation is used to analyze magnetic nanoantennas and is verified by our findings with full-wave simulations. We derive a formula that estimates the natural frequency associated to the magnetic resonance of a circular plasmonic cluster with an arbitrary number of plasmonic nanospheres. Lastly, we classify clusters based on their quality factor and their ability to enhance the magnetic field and discuss the surface area with strong magnetic field provided by the plasmonic cluster.

DOI: 10.1103/PhysRevApplied.8.024033

I. INTRODUCTION

The magnetic interaction of light and matter is weaker than its electric counterpart at optical frequencies [1-5]. Because of this reason, achieving resonant magnetism in the optical regime has become the focus of attention in the physics and engineering communities [6-12]. The overall magnetism of a metamaterial (i.e., relative magnetic permeability different from unity) is tailored by engineering the artificial magnetic resonance in “meta-atoms” [13,14]. For instance, engineered metamaterials with magnetic response were demonstrated in Refs. [15-19]. Split-ring resonators (SRRs) are the most explored building blocks in engineering artificial magnetism at the microwave spectrum [20-23]. Moreover, in Refs. [24-27], it is shown that by tuning the dimensions of SRRs and by implementing a precise fabrication, the magnetic resonance at near-infrared and visible frequencies is achievable. In Refs. [28-31], using effective-medium theory, a three-dimensional collection of polaritonic, nonmagnetic spheres have been shown to produce negative permeability at terahertz and infrared frequencies. Furthermore, other structures such as the spherical constellations [32-36], composite medium made of arrays of dielectric spheres [37], or a periodic lattice of clusters comprising four silver plasmonic dimers [38] have been used to make metamaterial constituents which provide magnetic polarization in the visible and infrared spectrums.

Circular clusters made of plasmonic nanoparticles, which are of interest in this paper, not only can provide engineered negative permeability [39], but they can also be used to achieve Fano resonances [40-44], and as magnetic nanoprobes, enhance light-matter interaction [45,46] at optical frequencies.

Conventionally, plasmonic nanoantennas are utilized at optical frequencies because of their exotic property of enhancing the electric field [47]. However, here we relate to the important quest to enhance magnetic field instead of the electric field [48]. In this paper, inspired by these referenced studies, we focus on the specific category of plasmonic nanoantennas made of circular clusters and focus on the enhancement of magnetic field. We refer to these antennas as magnetic nanoantennas or magnetic nanoprobes in the rest of the paper. Magnetic nanoantennas can be useful for enhancing quantum-magnetic transitions in molecules, which are, in general, overshadowed by the electric ones [45,46,49-52]. The magnetic resonance in a cluster of plasmonic nanoparticles corresponds to a dark resonance, i.e., a resonance that is neither easy to excite nor easy to measure its scattered field, which have higher quality factor and narrower bandwidth than the bright electric resonance counterpart. In Ref. [51], a cluster of six silver nanospheres has been studied under an azimuthally polarized beam, which selectively excites the dark resonance of the cluster with attention to the magnetic field enhancement and to the local field admittance of the scattered near field. However, to the best of the authors’ knowledge, the exploration of the best clusters for magnetic field enhancement and the amount or surface area where such enhancement occurs is still lacking in the literature as well as a simple analytic formula that estimates the magnetic resonance frequency given here.

Hence, in this paper, we investigate the properties of different clusters of gold nanospheres, such as dimers,
trimers, tetravers, pentamers, hexamers, and octamers in a host dielectric medium (e.g., glass or a general solution). Plasmonic gold nanoparticles have applications in medical diagnostics [53,54], sensing [55–58], and imaging [59] in the optical spectrum because of moderately low loss compared to several other metals, and also they are one of the least reactive chemical elements. Our goal is to investigate how magnetic nanoantennas made of gold are able to create a magnetic dominant region in which the magnetic field is enhanced over a certain surface area where the electric field ideally vanishes. Specific figures of merit (the magnetic field enhancement and the normalized local field admittance) are defined and used for that purpose as explained in the next section.

II. STATEMENT OF THE PROBLEM

We consider clusters of gold nanospheres with various numbers of elements, relative permittivity $\varepsilon_m$ in a host medium with relative permittivity $\varepsilon_h$ shown in Fig. 1. For the purpose of stressing the capabilities of magnetic nanoantennas in boosting quantum-magnetic transitions in matter that are overshadowed by electrical ones, we define two figures of merit [51]: the magnetic field enhancement and the magnetic-to-electric-field ratio, i.e., the normalized local field admittance:

$$F_H = \frac{|H'(r)|}{|H'(r)|},$$

$$F_Y = \eta \frac{|H'(r)|}{|E'(r)|}.$$

Here, $|H'(r)|$ and $|E'(r)|$ are the magnitude of total magnetic and electric fields at position $r$, and $|H'(r)|$ is the magnitude of incident magnetic field at the same position, respectively. The magnetic field enhancement ($F_H$) indicates the ability of a magnetic nanoantenna to enhance the magnetic field with respect to the incident one. The normalized magnetic-to-electric-field ratio, i.e., the normalized local field admittance ($F_Y$), shows the ability of a magnetic nanoantenna to enhance the magnetic field relative to the total electric field normalized to the field impedance $\eta = \sqrt{\varepsilon_0/\mu_0}$ of a plane wave in the same host medium. Large local field admittance in a region shows the magnetic nanoantenna succeeds in generating a magnetic dominant region, and $F_Y > 1$ means that the magnetic-to-electric-field ratio is larger than that of a plane wave.

Scattering from magnetic nanoantennas is characterized using two types of illuminations: (i) a single plane wave and (ii) two counterpropagating plane waves with antisymmetric electric field with respect to the cluster symmetry plane ($y$-$z$ plane in Fig. 1) to have a vanishing electric field at the center where a maximum of incident magnetic field occurs. The symmetry of illumination can be further augmented using azimuthally polarized beams with longitudinal magnetic field on their axis [51,60,61] whose field disposition matches perfectly to the magnetic mode of the clusters characterized by the rotationally symmetrical circulating electric dipole moments about the cluster center. The two counterpropagating plane-wave excitations are preferred over azimuthally polarized-beam illumination solely due to their instant availability and relaxed computation resource requirements in commercial electromagnetic full-wave-simulation software packages.

The cluster types used in our investigation, namely, dimers, trimers, tetravers, pentamers, hexamers, and octamers, are illustrated in Fig. 1. In all structures, the diameter of the nanospheres is $d$, and the gap between them is $g$. For a cluster made of $N$ spheres, each sphere is centered at the corners of a regular polygon whose circumscribed circle has radius $r_c$ called here the cluster radius given by

$$r_c = \frac{d + g}{2 \sin(\pi/N)}.$$

In this paper, the monochromatic time harmonic convention $\exp(-i\omega t)$ is implicitly assumed, and the notation is suppressed. In all equations, bold fonts are used for vector quantities in phasor domain, and a bar under a bold font is used for dyadic quantities. Unit vectors are bold with a hat on top.

III. ANALYTIC FORMULATION

We apply the single-dipole-approximation (SDA) method to model clusters of nanoparticles [62,63]. It means that we model each nanoparticle with a single electric dipole moment $p$. The electric dipole moment of the $n$th nanoparticle at location $r_n$ is found by

![FIG. 1. Plasmonic clusters with varying number of nanospheres: (a) dimer, (b) trimer, (c) tetramer, (d) pentamer, (e) hexamer, and (f) octamer. $r_c$ is the cluster radius, and $g$ is the gap between nanoparticles with diameter $d$.](Image)
\[
\mathbf{p}_n = \alpha \mathbf{E}^\text{loc}(\mathbf{r}_n),
\]
where \(\alpha\) is the electric polarizability of the nanosphere assumed to be isotropic, and \(\mathbf{E}^\text{loc}(\mathbf{r}_n)\) is the local electric field at the \(n\)th nanosphere’s location, which is the summation of the incident field and the field scattered by all the other nanospheres of the cluster. Here, the electric polarizability of the nanospheres is given by its Clausius-Mossotti expression with the correction term that accounts for the radiation [62]

\[
\alpha = \varepsilon_0 \varepsilon_h d^3 \left( \frac{2 \varepsilon_m + 2 \varepsilon_h - i (kd)^3}{\varepsilon_m - \varepsilon_h} \right)^{-1},
\]

where \(\varepsilon_h\) is the relative permittivity of the host medium, \(\varepsilon_0\) is the vacuum permittivity, \(d\) is the diameter of the nanosphere, \(k = k_0 \sqrt{\varepsilon_h}\) is the host medium wave number, and \(k_0\) is the wave number in vacuum. The gold nanosphere is described by its relative permittivity \(\varepsilon_m\) given here by the Drude model as

\[
\varepsilon_m = \varepsilon_\infty \left[ 1 - \frac{\omega_p^2}{\omega (\omega + i \gamma)} \right],
\]

where \(\varepsilon_\infty\) is the high-frequency fitting parameter, \(\omega_p\) is the plasma frequency, and \(\gamma\) is the damping factor. For gold, we assume \(\varepsilon_\infty = 9.5\), \(\omega_p = 4.4124 \times 10^{15}\) rad/s, and \(\gamma = 1.05 \times 10^{14}\) rad/s [64].

The total electromagnetic fields at an arbitrary observation point \(\mathbf{r}_{\text{obs}}\) is given by [65]

\[
\mathbf{E}'(\mathbf{r}_{\text{obs}}) = \mathbf{E}^\text{inc}(\mathbf{r}_{\text{obs}}) + \sum_{n=1}^{N} \mathbf{G}^\text{loc}(\mathbf{r}_{\text{obs}}, \mathbf{r}_n) \cdot \mathbf{p}_n,
\]

\[
\mathbf{H}'(\mathbf{r}_{\text{obs}}) = \mathbf{H}^\text{inc}(\mathbf{r}_{\text{obs}}) + \sum_{n=1}^{N} \frac{ck^2 e^{ikr}}{4\pi r} \left( 1 - \frac{1}{ikr} \right) (\mathbf{r}_{\text{obs}} \times \mathbf{p}_n),
\]

where \(c\) is the speed of light in the host medium, and \(\mathbf{G}^\text{loc}(\mathbf{r}_{\text{obs}}, \mathbf{r}_n)\) is the dyadic Green’s function defined as

\[
\mathbf{G}(\mathbf{r}_{\text{obs}}, \mathbf{r}_n) = \frac{e^{ikr}}{4\pi\varepsilon_0\varepsilon_h} \left[ \left( \frac{k^2}{r^3} + \frac{ik}{r^2} - \frac{1}{r} \right) \mathbf{I} - \left( \frac{k^2}{r^3} + \frac{3ik}{r^2} - \frac{3}{r} \right) \mathbf{r} \mathbf{r} \right].
\]

In Eqs. (8) and (9), \(r = |\mathbf{r}|\) with \(\mathbf{r} = \mathbf{r}_{\text{obs}} - \mathbf{r}_n\), where \(\mathbf{r}_n\) is the source dipole location, and \(\mathbf{I}\) is the \(3 \times 3\) identity dyad.

The overall electric dipole and overall magnetic dipole moments of a cluster of \(N\) nanospheres with the cluster center at the origin are defined as [44]

\[
\mathbf{p} = \sum_{n=1}^{N} \mathbf{p}_n, \quad \mathbf{m} = -\frac{i\omega}{2} \sum_{n=1}^{N} \mathbf{r}_n \times \mathbf{p}_n,
\]

respectively.

According to Eqs. (7) and (8), to calculate the electric and magnetic fields, the induced electric dipole for each sphere needs to be found. In doing so, we need to construct and solve a linear system of equations in terms of the induced dipole moments and the external excitation field. The local electric field \(\mathbf{E}^\text{loc}(\mathbf{r}_n)\) is given by

\[
\mathbf{E}^\text{loc}(\mathbf{r}_n) = \mathbf{E}'(\mathbf{r}_n) + \frac{\sum_{m=1}^{N} \mathbf{G}(\mathbf{r}_n, \mathbf{r}_m) \cdot \mathbf{p}_m}{\sum_{m=1}^{N} \mathbf{p}_m}. \quad (11)
\]

By writing Eqs. (4) and (11) for \(n = 1, \ldots, N\), we can construct the linear system

\[
[A] \begin{bmatrix} \mathbf{p}_1 \\ \vdots \\ \mathbf{p}_N \end{bmatrix} = \begin{bmatrix} \alpha \mathbf{E}'(\mathbf{r}_1) \\ \vdots \\ \alpha \mathbf{E}'(\mathbf{r}_N) \end{bmatrix}, \quad (12)
\]

where \([A]\) is a \(3N \times 3N\) matrix made of \(3 \times 3\) sub-blocks \(\Delta_{mn}\), with \(n, m \in \{1, \ldots, N\}\) given by

\[
\Delta_{mn} = \left\{ \begin{array}{ll} 1 & m = n, \\ -\alpha \mathbf{G}(\mathbf{r}_n, \mathbf{r}_m) & m \neq n. \end{array} \right. \quad (13)
\]

As mentioned earlier, one of the goals of this paper is to sort the magnetic nanoantennas based on their quality factor associated to the magnetic resonance, which depends on both material and radiation losses that are accounted for by using the dynamic Green’s function [Eq. (9)]. One way to calculate the quality factor of each cluster is to find the cluster natural frequencies. To do so, we need to solve Eq. (12) for \(\omega\) when \(\mathbf{E}_i = 0\). One has nontrivial solutions to such a system only when

\[
\det[A] = 0. \quad (14)
\]

In general, solving Eq. (14) for \(\omega\) gives complex frequencies. One way to calculate the natural frequencies is to solve Eq. (14) numerically. Under certain approximations, one can also obtain analytical formulas of resonance frequencies. Similarly, in Ref. [66], the authors applied SDA to calculate the natural frequency of a dimer of particles made of metallic nanoshell and dielectric core for symmetric and antisymmetric conditions. In the next section, we study only the rotationally symmetric magnetic resonance case by simplifying Eq. (14) to a scalar equation, and we provide a simple approximate formula to estimate the natural frequency and quality factor of clusters with an arbitrary number of metal nanospheres.
FIG. 2. The magnetic resonance has a symmetric disposition of electric dipoles along $\hat{\phi}$ producing a strong magnetic field at the center.

A. Magnetic resonance frequency

Because of the rotational symmetry associated to a magnetic resonance in a cluster, each nanosphere induces a circulating displacement current resulting in the rotationally symmetric electric-dipole-moment disposition shown in Fig. 2. This resonance generates the overall longitudinal magnetic dipole moment of the cluster provided in Eq. (10). Therefore, because of symmetry, we reduce the system in Eq. (12) made of $3N$ equations to a single scalar equation. This is because each sphere has identical induced electric-dipole-moment amplitude $p_{\varphi}$ and polarized along the $\hat{\phi}$ direction as $p_{\varphi} = p_{\varphi} \hat{\phi}$. Thus, the local electric field at the $n$th nanosphere is written as $E^{(n)}(r_n) = (p_{\varphi}/\alpha) \hat{\phi}_n$. So, Eq. (11) is written as

$$
(p_{\varphi}/\alpha) \hat{\phi}_n - \sum_{m=1 \neq n}^{N} G(r_n, r_m) \cdot p_{\varphi} \hat{\phi}_m = E'(r_n). \tag{15}
$$

Natural frequencies are the solution of Eq. (15) without an excitation wave ($E' = 0$). By dividing all vector terms in Eq. (15) by $\alpha^{-1} p_{\varphi}$ and projecting the vectors in Eq. (15) on $\hat{\phi}_n$, we obtain

$$
1 - \sum_{m=1 \neq n}^{N} \alpha \hat{\phi}_n \cdot G(r_n, r_m) \cdot \hat{\phi}_m = 0. \tag{16}
$$

In Sec. IV, the complex natural frequency solution of Eq. (16) is also calculated numerically.

A closed-form formula for the complex natural frequency is obtained by applying some approximations. The first assumption is that both nanospheres and their mutual distances are subwavelength. Therefore, it is possible to replace the dynamic terms in both the polarizability and the Green’s functions by the electrostatic ones, which means the quasistatic solution is equivalent to the limiting case where $k \to 0$ in Eqs. (5) and (9) but keeping the correct frequency-dependent dielectric constant $\varepsilon_m$ in Eq. (6). The polarizability reduces to the Clausius-Mossotti relation [62]

$$
\alpha \approx \alpha_0 = \frac{1}{2} \pi \varepsilon_0 \varepsilon_h \rho_d \left( \frac{\varepsilon_m - \varepsilon_h}{\varepsilon_m + 2\varepsilon_h} \right). \tag{17}
$$

The Green’s function is approximated in the static regime as [63]

$$
G(r_n, r_m) \approx \frac{1}{4\pi \varepsilon_0 \varepsilon_h} \left( -\frac{1}{r_{nm}} \mathbf{I} + \frac{3}{r_{nm}^3} \mathbf{r}_{nm} \mathbf{r}_{nm} \right). \tag{18}
$$

By substituting Eqs. (17) and (18) in Eq. (16), after some algebraic manipulation, Eq. (16) is reduced to

$$
e_m = 1 + \frac{2L}{1-L} \varepsilon_h, \tag{19}
$$

where $L = (2r_c/d)^3/T$ and

$$
T = \sum_{n=1 \neq n}^{N} \left\{ \frac{r_c}{r_{nm}} \frac{3}{D_{nm}} \left[ \frac{3}{D_{nn}} - 1 \right] \sin \varphi_n \sin \varphi_m - \frac{3B_{nm}}{D_{nm}} \sin(\varphi_n + \varphi_m) + \frac{3C_{nm}}{D_{nm}} - 1 \right] \cos \varphi_n \cos \varphi_m \right\}. \tag{20}
$$

In Eq. (20), $A_{nm}$, $B_{nm}$, $C_{nm}$, and $D_{nm}$ are the coefficients which relate the $n$th nanosphere to the $m$th one and are defined as

$$
A_{nm} = (\cos \varphi_n - \cos \varphi_m)^2,
B_{nm} = (\cos \varphi_n - \cos \varphi_m)(\sin \varphi_n - \sin \varphi_m),
C_{nm} = (\sin \varphi_n - \sin \varphi_m)^2,
D_{nm} = A_{nm} + C_{nm}. \tag{21}
$$

We are looking for a complex natural angular frequency as $\omega = \omega' + i\omega''$. Substituting the Drude model for metal permittivity Eq. (6) into Eq. (19) and solving for $\omega$ leads to the closed-form formula for complex natural angular frequency as

$$
\omega' \approx \sqrt{\frac{\omega_p^2 \varepsilon_h}{\varepsilon_{\infty} - \frac{1+2L}{1-L} \varepsilon_h} - \frac{\gamma^2}{4}}, \quad \omega'' \approx \frac{\gamma}{2} \tag{22}
$$

Note that Eq. (22) reduces to the natural frequency reported in Ref. [67] that is obtained for the specific case with $N = 4$ nanospheres.

It is important to note that with the quasistatic approximation, we neglect the radiation damping; therefore, material loss (based on the Drude model) is the only remaining loss when evaluating these formulas. Since the natural frequency is complex, one can define the quality factor as [68]

$$
Q = \frac{\omega'}{2\omega''}. \tag{23}
$$

A system with high quality factor provides a sharp resonance and strong overall magnetic dipole moment.
According to the derived approximate formula for the natural frequency [Eq. (22)], we find an approximate formula for the quality factor of clusters with an arbitrary number of nanospheres as

$$ Q \approx \sqrt{\frac{(\omega_p/\gamma)^2}{\epsilon_\infty - \frac{1}{1+\frac{4}{\pi L}} \epsilon_h}} - \frac{1}{4}. $$

where the number of nanospheres ($N$) is accounted for in the $L$ parameter. In the next section, we report the approximate (only material loss is considered) quality factor and natural frequency of the clusters of Fig. 1 with a comparison to the numerically found ones (both material loss and radiation loss are considered).

**IV. RESULTS AND DISCUSSION**

We address the effect of cluster parameters on the resonance frequency and on the two figures of merit $F_H$ and $F_Y$ introduced in Sec. II. The structures we study include dimers, trimers, tetramers, pentamers, hexamers, and octamers embedded in a host medium with relative permittivity $\epsilon_h = 2.25$ (it can represent the permittivity of an environment consisting of a glass substrate and a solution).

In the following, the magnetic resonance frequency is calculated using two distinct methods: (i) by solving Eq. (14) for complex frequency, in two ways: numerically, when the electric polarizability in Eq. (5) is used [denoted $F_{\text{III}}$ in Table I] and by using the closed-formula solution obtained via the approximate analytical method introduced in Sec. III [denoted as in Table I], and (ii) by finding the purely real resonance frequency defined as the frequency that renders the magnetic-field-enhancement $F_H$ maximum at the cluster center under time harmonic excitation [resonance denoted by $f_r(\text{III})$ in Table I]. In this latter case, we assume the cluster illuminated by a single plane wave with electric field polarized along the $y$ axis and magnetic field polarized along $z$, as shown in Fig. 3(a).

In the reported comparisons of resonance frequencies calculated by different methods, we consider the constant sphere diameter and gap in the clusters of Fig. 1 as 50 and 5 nm, respectively. In the fifth column of Table I, we report $F_H$ at the resonance frequency under single-plane-wave excitation. Finally, the quality factor $Q$ based on the natural resonance frequency calculated numerically $f_r(\text{I})$ and with the approximate formula $f_r(\text{II})$ as in Eq. (22) are reported in the sixth and seventh columns of Table I, respectively. Furthermore, to provide a clear comparison among the methods, we define the percentage error in the real part of frequencies calculated numerically [$f_r(\text{I})$] and analytically (by approximate formula) [$f_r(\text{II})$] relative to $f_r(\text{III})$ as

$$ |\epsilon_r|\% = \left| \frac{f_r(\text{I or II}) - f_r(\text{III})}{f_r(\text{III})} \right| \times 100. $$

For the clusters reported in Table I, the relative error for the natural frequency calculated numerically [$f_r(\text{I})$] and approximately [$f_r(\text{II})$] is always less than 1% and 10%, respectively. The very good agreement between $f_r(\text{I})$ and

### TABLE I. Comparison of natural frequencies found with different methods and the quality factor of different clusters of gold nanospheres when $d = 50$ nm and $g = 5$ nm.

| Cluster $N$ | $f_r(\text{I})$ THz $f_r(\text{II})$ THz $f_r(\text{III})$ THz $F_H$ $Q(\text{I})$ $Q(\text{II})$ |
|-------------|-----------------|-----------------|-----------------|---------|--------|--------|
| 2           | 566-i10         | 564-i8          | 566             | 9       | 28.6   | 33.8   |
|             | $|\epsilon_r| = 0.1\%$ | $|\epsilon_r| = 0.2\%$ |                 |         |        |        |
| 3           | 538-i10         | 540-i8          | 537             | 6.7     | 26.6   | 32.3   |
|             | $|\epsilon_r| = 0.2\%$ | $|\epsilon_r| = 0.5\%$ |                 |         |        |        |
| 4           | 513-i11         | 523-i8          | 511             | 7.1     | 22.6   | 31.3   |
|             | $|\epsilon_r| = 0.4\%$ | $|\epsilon_r| = 2\%$ |                 |         |        |        |
| 5           | 494-i13         | 513-i8          | 492             | 6.5     | 18.9   | 30.7   |
|             | $|\epsilon_r| = 0.3\%$ | $|\epsilon_r| = 4\%$ |                 |         |        |        |
| 6           | 479-i15         | 506-i8          | 477             | 5.8     | 15.6   | 30.3   |
|             | $|\epsilon_r| = 0.3\%$ | $|\epsilon_r| = 6\%$ |                 |         |        |        |
| 8           | 456-i21         | 500-i8          | 452             | 4.5     | 10.8   | 29.9   |
|             | $|\epsilon_r| = 1\%$ | $|\epsilon_r| = 10\%$ |                 |         |        |        |
The quality factor calculated numerically $(Q(III))$ is consistent with the high quality factor $Q(I)$, which means that the resonant mode is clearly excited and well defined even by single-plane-wave excitation (we recall that to purely excite the magnetic resonance, two counter-propagating plane waves need to be used, otherwise, the single plane wave excites also other cluster modes). Note also that our approximate formula Eq. (22) provides a good estimate of resonance frequency. Moreover, by looking at the magnetic field enhancement $F_H$ and the quality factor $Q$ calculated numerically and approximately, two things can be inferred: (i) our approximate formula for the quality factor works as a tool to show how strong the magnetic nanoantenna enhances the magnetic field, and (ii) as the number of elements in a cluster increases, the magnetic field enhancement at the cluster center and the quality factor decrease.

As shown in Appendix A, for a cluster with number of nanospheres $N$, assuming the resonance is made of $N$-induced electric dipole moments perfectly polarized along $\phi$ as shown in Fig. 2, the ratio of scattered power to the absorbed power is proportional to $P^{\text{scat}}/P^{\text{abs}} \propto N/N^2(\pi/N)$. This means that when $N$ increases, for the magnetic resonance in these clusters, the loss due to scattering (as a magnetic dipole) is larger than the loss in the nanospheres. Hence, when the cluster is enlarged by adding more nanospheres (whereas the diameter of each nanosphere and the gap between them is fixed), the quality factor calculated numerically $(Q(I))$ (which is based on both material and radiation loss) drops more rapidly than the one calculated approximately $(Q(II))$ (which is based only on material loss).

Next, in Fig. 4 we explore the effect of the nanospheres’ diameter $d$ and gap spacing $g$ on the magnetic field enhancement $F_H$ (evaluated at the cluster center at its resonance frequency) for the clusters shown in Fig. 1.

We excite each cluster with a single plane wave polarized along the $y$ direction as shown in Fig. 3(a). For each cluster, the nanosphere diameter varies from 5 to 90 nm, and the gap varies from 1 to 20 nm. Note that the resonance frequency is not constant when varying $d$ and $g$; therefore, in Fig. 4, we superimpose some of the isofrequency contours denoting the resonance frequency at which $F_H$ is given.

The results shown in Fig. 4 verify the important conclusion that as the cluster’s number of nanospheres increases, the magnetic field enhancement at its center decreases. Furthermore, according to Fig. 4, for a given $N$ and $g$, the magnetic field enhancement at the cluster center increases by increasing the nanosphere diameter until it reaches a maximum for an optimum diameter, and if the diameter is further increased, $F_H$ decreases. Moreover, according to Fig. 4, for a given sphere diameter, $F_H$ decreases by increasing the gap space between the two adjacent nanospheres.

Figure 5 shows the normalized local field admittance $F_Y$ at the center of a hexamer evaluated at its resonance frequency, which depends on the varying parameters $d$ and $g$. According to Fig. 5, the effect of the diameter and gap on the local field admittance at the center of the

FIG. 4. Magnetic field enhancement $F_H$ at the center of each cluster at its resonance frequency for the (a) dimer, (b) trimer, (c) tetramer, (d) pentamer, (e) hexamer, and (f) octamer. The nanospheres’ diameter $d$ and gap $g$ vary between 5 and 90 nm and between 1 and 20 nm, respectively. Each cluster is excited by a single plane wave polarized along the $y$ axis.
About resonating clusters at a certain resonance frequency (here we choose 480 THz, since most of the clusters have resonance at this frequency). Figure 6 represents the magnetic field enhancement ($F_H$) at the cluster center versus the radius of the cluster ($r_c$) for clusters with various number of nanospheres which resonate at 480 THz. It is worth mentioning that for each cluster with a specific number of nanospheres ($N$) resonating at 480 THz, there is an optimum cluster radius at which the magnetic field enhancement is maximized. Moreover, when the number of nanospheres in a cluster increases, the optimum radius of the cluster increases, and its corresponding magnetic field enhancement reduces.

As mentioned earlier, to create a magnetic dominant region, one needs to establish symmetry conditions on both the cluster and the incident field such that the net electric response of the cluster is suppressed. The clusters we introduced so far are rotationally symmetric and host a strong magnetic dipolar mode. Moreover, by applying two counterpropagating plane waves with antisymmetric electric field distribution with respect to cluster symmetry plane ($y$-$z$ plane in our structure) as shown in Fig. 3(b), the external electric field vanishes. In addition, due to the cluster’s rotational symmetry, no net electric field is created at the cluster center. This excitation method leads to strong electric field in the gap region between nanospheres and strong magnetic field at the cluster center. Our findings (for brevity, the details are not shown here) show that for the clusters studied in this paper, the magnetic field enhancement defined in Eq. (1) does not depend on the excitation type, i.e., a single plane wave versus two counterpropagating plane waves as in Fig. 3(b). With two counterpropagating plane-wave illuminations, for each cluster, we study the behavior of magnetic field enhancement with respect to frequency based on both the analytical formulation using SDA and full-wave simulations. Full-wave simulations are performed by the frequency-domain finite-element method implemented in the commercial software CST Microwave Studio by Computer Simulation Technology AG. Figure 7 shows the magnetic field enhancement ($F_H$) versus
frequency for the clusters shown in Fig. 1 at the cluster center by using SDA [Fig. 7(a)] and full-wave simulations [Fig. 7(b)] when the diameter of each nanosphere and the gap spacing are kept constant as 50 and 5 nm, respectively, and the clusters are excited by two counterpropagating plane waves as shown in Fig. 3(b).

Figure 7 confirms that both analysis methods lead to the same conclusion that enlarging the cluster by adding more nanospheres results in reducing the magnetic field enhancement at the cluster center, redshifting the resonance frequency, and increasing the resonance line width. There is a slight difference between the resonance frequencies obtained with full-wave simulation and those obtained with SDA (5% in the dimer case and increasing to 15% in the octamer case). The SDA calculations also overestimate the magnetic field enhancement by 8% in the trimer case and up to 50% in the octamer case. The difference between the results obtained with these two methods grows as the number of nanospheres increases. This difference originates from the fact that in SDA we neglect the magnetic dipole moment of each nanosphere and all higher-order multipoles, and by adding more nanospheres in a cluster, the effect is cumulative. To elaborate on this issue, we study also the absorption [Figs. 8(a) and 8(b)], scattering [Figs. 8(c) and 8(d)], and extinction [Figs. 8(e) and 8(f)] cross sections of the clusters shown in Fig. 1 based on the equations given in Appendix B. These cross sections are evaluated using both the SDA [Figs. 8(a), 8(c), and 8(e)] and full-wave simulations [Figs. 8(b), 8(d), and 8(f)] assuming the diameter of each nanosphere and the gap spacing to be constant as 50 and 5 nm, respectively. In these simulations, the clusters are excited by two counterpropagating plane waves as shown in Fig. 3(b). In each plot, the cross sections are normalized to the geometrical cross section of a single nanosphere $\sigma_g = \pi(d/2)^2$. At the magnetic resonance, each cluster exhibits a peak in the absorption and scattering and, hence, in the extinction cross section. Moreover, according to the cross sections specifically obtained by full-wave simulations, we observe that adding more nanospheres in a cluster leads to a larger cross section. Furthermore, since the scattering cross section is proportional to the total magnetic dipole moment of the cluster, as the number of nanospheres increases, stronger magnetic dipole moment is achieved. We use two counterpropagating plane-wave excitations since it facilitates the simulation burden, and it makes the electric dipole moment vanish. Higher-order multipoles like the electric quadrupole are still present, though weak compared to the magnetic dipole since each of these simulations is carried out at the magnetic resonance frequency. The effects of the electric quadrupole are visible at higher frequencies in the simulations, analogous to what is shown in detail in Figs. 3–5 of Ref. [33] for spherical clusters. Note that higher-order multipoles are not visible in the scattering cross section calculated with the SDA since it is evaluated in an approximate way [Eq. (A5) by considering only the scattering due to a magnetic dipole].

The results also confirm the trend already observed in Fig. 7: as we increase the number of nanospheres in the cluster, the results of the SDA diverge from those of full-wave simulations.

Although the magnetic field enhancement reduces at the cluster center as the cluster expands, to elaborate on the possible advantages of large clusters, we provide the magnetic-field-enhancement profile in the x-y plane for different clusters at their resonance frequency. Figures 9(a)–9(f) represent the magnetic-field-enhancement ($F_H$) profile in logarithmic scale [as 10log$_{10}(F_H)$] the color legend is saturated for values more than 10.4] calculated using SDA analysis for the clusters represented in Fig. 1 when the nanospheres’ diameter and gap between them are kept constant as 50 and 5 nm, respectively, and the clusters are excited by two counterpropagating plane waves as shown in Fig. 3(b) at their resonance frequency obtained from Fig. 7(a). According to Fig. 9, the magnetic field enhancement at the gap space between adjacent nanospheres is
always strong, and it is slightly stronger near the nanospheres rather than at the cluster center. Moreover, as the number of nanospheres in the cluster increases, the area with strong magnetic field becomes wider. According to Table I, the quality factor decreases with size, which results in the reduction of the magnetic field enhancement at the cluster center. Therefore, there is always a trade-off between getting the maximum magnetic field enhancement at a certain point such as the center of a cluster and having strong magnetic field over a wide area inside the cluster. The plots in Fig. 9 show also a strong magnetic field enhancement outside the cluster with the shape of two vertical bands (dark red bands with extraordinarily large $F_H$). This phenomenon is due to the specific excitation used in Fig. 9 with two counterpropagating plane waves shown in Fig. 3(b) that form a standing wave pattern of the incident magnetic field. There are two nulls with half a wavelength distance and result in locally huge magnetic field enhancement [51].

Furthermore, to demonstrate the advantage of the symmetric excitation method [as shown in Fig. 3(b)] and to create the magnetic dominant region with low electric field, we investigate the local field admittance ($F_y$) profile in the $x$-$y$ plane when the clusters are excited with two counterpropagating plane waves as in Fig. 3(b). Figures 10(a)–10(f) show the local field admittance profile in logarithmic scale (the color legend is saturated for the values more than 4) calculated using the SDA method for clusters shown in Fig. 1 with fixed diameter and gap of 50 and 5 nm, respectively, while the clusters are excited with two counterpropagating plane waves with antisymmetric electric field with respect to the $y$-$z$ plane.

Because of the two counterpropagating plane-wave excitations shown in Fig. 3(b), the incident electric fields cancel out exactly at the center of the cluster, so a giant local field admittance is expected at the center of each cluster. Moreover, enlarging the clusters by adding more nanospheres leads to a wide area inside the cluster which possesses strong magnetic field (Fig. 9) and large field admittance (Fig. 10). In other words, there is a region of magnetic dominance. Furthermore, a comparison of Fig. 10(e) and Fig. 5 for the case of $d = 50$ nm and $g = 5$ nm clearly shows that the level of local field ADMITTANCE enhances with the number of nanospheres in the cluster, which results in a wider area with strong magnetic field enhancement.
admittance is much higher when the clusters are excited by a symmetric excitation.

V. CONCLUSION

We investigate clusters of gold nanospheres embedded in a host medium as magnetic nanoantennas to enhance the magnetic field at optical frequencies, and we show how the electric field can be reduced at the cluster center. We sort clusters based on their quality factor, the magnetic field enhancement, and their ability to create a wide magnetic dominant region. We apply SDA to calculate the total magnetic and electric fields at an arbitrary point while the clusters are illuminated by two different methods of excitation: (i) single plane wave and (ii) superposition of two counterpropagating plane waves to eliminate the electric field at the cluster center where the magnetic field is enhanced. We provide a formula to approximate the natural frequency of clusters and their quality factor with an arbitrary number of elements. We also calculate the natural frequency and quality factor numerically and demonstrate the validity range of our proposed formulation. We define two figures of merit to study the magnetic nanoantenna’s efficacy: (i) magnetic field enhancement ($F_H$), which shows the ability of the nanoantenna to enhance the magnetic field, and (ii) the local field admittance ($F_Y$), which shows how much the magnetic field is enhanced compared to the electric field. We verify our analytic results against full-wave simulations and show their consistency. The results indicate three facts about the clusters of nanospheres as magnetic nanoantennas: (i) Increasing the number of elements in a cluster leads to redshift in resonance frequency, decrease in quality factor, and the level of magnetic field enhancement at the center of the cluster, whereas the area of the magnetic dominant region increases. (ii) In each cluster, increasing the diameter of spheres first causes stronger enhancement, and there is an optimum diameter after which the enhancement reduces by further increasing the diameter. (iii) Increasing the gap spacing in a cluster causes less magnetic field enhancement at the cluster center. According to our study, clusters of gold nanospheres, because of their symmetry, when illuminated by a symmetric excitation are good magnetic nanoantennas with large magnetic-to-electric-field ratio at the center.

FIG. 10. The local field admittance ($F_Y$) profile of different clusters with $d = 50$ nm and $g = 5$ nm in logarithmic scale (the color legend is saturated for the values more than 4, i.e., for $F_Y > 10^4$) for the (a) dimer, (b) trimer, (c) tetramer, (d) pentamer, (e) hexamer, and (f) octamer. Each cluster is excited with two counterpropagating plane waves with antisymmetric electric field with respect to the $y$-$z$ plane. The results show a large magnetic-to-electric-field ratio around the center.
ACKNOWLEDGMENTS

The authors acknowledge support from the W. M. Keck Foundation, USA, and from the National Science Foundation, Grant No. NSF-SNM-1449397. The authors are grateful to CST Simulation Technology AG for letting them use the simulation tool CST Microwave Studio that is instrumental in this analysis.

APPENDIX A: RADIATED AND ABSORBED POWER BY A CLUSTER

We assume that for a resonating cluster with \( N \) nanospheres, the magnetic resonance has a symmetric disposition of electric dipoles as shown in Fig. 2. Hence, all the \( N \) electric dipoles are polarized along the \( \hat{y} \) direction and have equal magnitude \( p_\omega \). According to Refs. [69,70], the absorbed time-average power in the \( n \)th nanosphere of the cluster

\[
P_{\text{abs}} = \frac{\alpha \omega_0 \epsilon_0}{2} \left[ \frac{\text{Im}(\alpha)}{\epsilon_0 \epsilon_h} - \frac{k^3}{6 \pi \epsilon_0 \epsilon_h} \right] |p_\omega|^2 \quad (A1),
\]

where \( \alpha \) is the polarizability of each nanosphere given in Eq. (5). Moreover, since we assume that at resonance \( \epsilon_\omega \) is the radius of the cluster defined in Eq.(3). The power absorbed by the cluster to the absorbed power is written as

\[
P_{\text{abs}} = \frac{\alpha \omega_0 \epsilon_0}{2} \left[ \frac{\text{Im}(\alpha)}{\epsilon_0 \epsilon_h} + \frac{k^3}{6 \pi \epsilon_0 \epsilon_h} \right] |p_\omega|^2. \quad (A2)
\]

Because of the circular symmetry of the dipoles’ strength, the total cluster power lost due to absorption is

\[
P_{\text{abs}} = \sum_{n=1}^{N} P_{\text{abs}} = \frac{\alpha \omega_0 N}{2} \left[ \frac{\text{Im}(\alpha)}{\epsilon_0 \epsilon_h} + \frac{k^3}{6 \pi \epsilon_0 \epsilon_h} \right] |p_\omega|^2. \quad (A3)
\]

According to Eq. (10), and because of the symmetry of the magnetic resonance, the overall magnetic dipole moment of the cluster with \( N \) nanospheres when the cluster center is at the origin is

\[
\mathbf{m} = -\frac{i \omega}{2} r_c p_\omega N \hat{\mathbf{z}}, \quad (A4)
\]

where \( r_c \) is the radius of the cluster defined in Eq. (3). The scattered power by the cluster is evaluated as the power scattered by a magnetic dipole moment leading to [71]

\[
P_{\text{scat}} = \frac{\alpha \omega}{2} \frac{k^3 \mu_0^2}{\epsilon_0 \epsilon_h} \left( \frac{|\mathbf{m}|^2}{\eta^2} \right), \quad (A5)
\]

where we neglect the power contributions associated to higher-order multipoles. By substituting Eq. (A4) into Eq. (A5), the total scattered power by the cluster is

\[
P_{\text{scat}} = \frac{\alpha^5}{48\pi} k_0 \epsilon_0 \epsilon_h r_c^2 N^2 |p_\omega|^2. \quad (A6)
\]

If we assume that in a cluster with \( N \) nanospheres, the diameter of each nanosphere and the gap between them is kept constant when varying \( N \), by substituting the expression for the cluster radius in Eq. (3) into Eq. (A6), the ratio of scattered power by a cluster to the absorbed power is

\[
\frac{P_{\text{scat}}}{P_{\text{abs}}} = -\frac{N}{\sin^2(\pi/N)} \frac{\alpha^4}{96\pi} \frac{\epsilon_\omega}{\epsilon_0} \left| \text{Im}(\alpha) \right| + \frac{N^4}{6\pi \epsilon_0 \epsilon_h}. \quad (A7)
\]

Moreover, one can substitute the Drude model permittivity provided in Eq. (6) into the electric polarizability [Eq. (5)], and then the ratio of the scattered power by a cluster to the absorbed power is written as

\[
\frac{P_{\text{scat}}}{P_{\text{abs}}} = \frac{N}{\sin^2(\pi/N)} \frac{Z}{\eta^2}, \quad (A8)
\]

where

\[
Z = \frac{\alpha^3 \epsilon_\omega \epsilon_0 \mu_0^2 k \pi d^2 (d + g)^2}{576} \left\{ \frac{\omega^2 (\epsilon_\omega - \epsilon_h) - \epsilon_\omega \omega_p^2}{\gamma \epsilon_\omega \omega_p^2} \right\}. \quad (A9)
\]

APPENDIX B: ABSORPTION, SCATTERING, AND EXTINCTION CROSS SECTIONS

According to Ref. [72], the absorption, scattering, and extinction cross sections are defined as

\[
\sigma_{\text{abs}} = \sum_{n=1}^{N} \frac{P_{\text{abs}}}{S^n}, \quad \sigma_{\text{scat}} = \frac{P_{\text{scat}}}{S^n}, \quad (B1)
\]

and \( \sigma_{\text{ext}} = \sigma_{\text{abs}} + \sigma_{\text{scat}} \). Here, \( P_{\text{abs}} \) and \( P_{\text{scat}} \) are calculated using Eqs. (A1) and (A5), and \( S^n = (\eta/2) |\mathbf{H}^n|^2 \), where \( |\mathbf{H}^n| \) is the magnitude of the total incident magnetic field exciting the cluster metaparticle. Note that we are focusing on the magnetic dipole excitation of a cluster that is seen here as a single scattering and absorbing metaparticle. Indeed, scattering of a particle described by its magnetic dipole is proportional to \( |\mathbf{m}|^2 = |\mathbf{m}^{\text{eff}}|^2 \) (\( \epsilon_\omega \) is the effective magnetic polarizability of the cluster metaparticle) as shown in Eq. (A5), and its absorption is proportional to \( \text{Im}(\mathbf{m} \cdot \mathbf{H}^{\text{inc}}) \propto |\mathbf{H}^n|^2 \), where the asterisk denotes complex conjugation. This case is analogous to that of a particle described by its electric dipole absorption and scattering that is proportional to \( S^i = (1/2\eta) |\mathbf{E}^i|^2 \). Note that in our simulations where we use an excitation made of two counterpropagating plane waves [as shown in Fig. 3(b)], \( S^i \) defined above is equivalent to \( S^i = 4k_{\text{PW}}^2 \Re(\mathbf{E}^{\text{PW}} \times \mathbf{H}^{\text{PW}}) = 4(\eta/2)|\mathbf{H}^{\text{PW}}|^2 \), where \( k_{\text{PW}} \) is normalized wave vector of
either plane wave, and $\mathbf{E}^{PW}$ and $\mathbf{H}^{PW}$ are its electric and magnetic fields. For the case of two counterpropagating plane waves considered in Fig. 3(b), we have $\mathbf{H}^i = 2\mathbf{H}^{PW}$ and $\mathbf{E}^i = 0$ at the cluster center where we assume the equivalent magnetic dipole is located.


[41] B. Luk’yanchuk, N. I. Zheludev, S. A. Maier, N. J. Halas, P. Nordlander, H. Giessen, and C. T. Chong, The Fano resonance in plasmonic nanostructures and metamaterials, Nat. Mater. 9, 707 (2010).


[64] N. K. Grady, N. J. Halas, and P. Nordlander, Influence of dielectric function properties on the optical response of