Decay Rate of Magnetic Dipoles near Non–magnetic Nanostructures

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In this article, we propose a concise theoretical framework based on mixed field-susceptibilities to describe the decay of magnetic dipoles induced by non–magnetic nanostructures. This approach is first illustrated in simple cases in which analytical expressions of the decay rate can be obtained. We then show that a more refined numerical implementation of this formalism involving a volume discretization and the computation of a generalized propagator can predict the dynamics of magnetic dipoles in the vicinity of nanostructures of arbitrary geometries. We finally demonstrate the versatility of this numerical method by coupling it to an evolutionary optimization algorithm. In this way we predict a structure geometry which maximally promotes the decay of magnetic transitions with respect to electric emitters.

PACS numbers: 68.37.Uv Near-field scanning microscopy and spectroscopy
78.67-n Optical properties of low-dimensional materials and structures
73.20.Mf Collective excitations

I. INTRODUCTION

During the last two decades, the development of nanooptics has provided a wealth of strategies to tailor electric and magnetic fields down to the subwavelength scale. In particular, optical nano-antennas have allowed to modify the intensity, dynamics or directionality of light emission from fluorophores placed in the near-field of nano-objects using concepts from the radiofrequency domain. These studies have been performed nearly exclusively on fluorophores supporting electric dipole (ED) transitions, the latter being $\sim a_0/\lambda_0 \sim 10^{-4} - 10^5$ larger than their magnetic dipole (MD) counterpart in the optical frequency range ($a_0$ being the Bohr radius and $\lambda_0$ the transition wavelength). Recently, delicate experiments have addressed light emission from rare earth doped emitters supporting both strong MD and ED transitions. Three-dimensional maps of the luminescence of Eu$^{3+}$ doped nanocrystals scanned in the near-field of gold stripes have revealed variations in the relative intensities of ED and MD transitions. In these experiments, the fluorescence intensity, photon statistics and branching ratios are directly related to the decay rates of the ED or MD radiative transitions, the latter being ultimately connected to the electric or magnetic part of the local density of electromagnetic states (EM-LDOS). Independently of the nature of the transition, the alteration of the EM-LDOS by a nanostructure arises from the back action of the electric or magnetic field on the transition dipole. The confinement of the magnetic field around non-magnetic nano-objects arises from the spatial variations of the electric near-field in the immediate proximity of a nanostructure. When the surface is illuminated by a plane wave or an evanescent surface wave, both experimental data and numerical simulations reveal spatial modulations in the electric and magnetic near-field intensities. For example, the magnetic field intensity recorded above subwavelength sized dielectric particles, excited by a p–polarized surface wave, has a strong and dark contrast while a completely opposite behavior is observed for the electric field intensity. If now, the nanostructure is illuminated, no longer by a plane wave but by a dipole source, the response fields (electric or magnetic) are different and shape the decay rate and the corresponding dipolar luminescence. From a mathematical point of view, the magnetic near-field can be described by a set of mixed field–susceptibilities capable of connecting an electrical polarization, oscillating at an optical frequency $\omega_0$, to a magnetic field vector oscillating at the same frequency. In fact, these field–susceptibilities are a generalized form of the usual Green dyadic tensor. Historically, they were introduced by G.S. Agarwal to describe energy transfers in the presence of dielectric or metallic planar surfaces. Mixed field–susceptibilities can be used to evaluate the optical magnetic near-field or the optical response of nano-structures possessing an intrinsic magnetic polarizability, like metallic rings or split-rings. In recent works, they have been used to separately study the magnetic and electric part of the LDOS close to a surface and for the calculation of the EM-LDOS in proximity of periodic arrays of magneto-electric point scatterers.

In this article, we first extend Agarwal’s theory by presenting a new analytical scheme yielding the total decay rate of a MD transition $\Gamma_{m}$ in terms of mixed electric–
The two possible kinds of coupling between a magnetic dipole transition and a subwavelength sized sphere are schematized in figure 1-a and b. The first one is the direct magnetic coupling such as the one involved in the presence of artificial left-handed materials, i.e. materials with simultaneously negative permeability and permittivity. We address exclusively the first situation and therefore assume that $\mu(\omega_0) = 1$ at all wavelengths. We consider the geometry depicted in figure 1. The electric and magnetic fields generated at $r$ by a magnetic dipole $m(\omega_0)$ located at $r_0$ are defined by:

$$E_0(r, \omega_0) = i k_0 \nabla_r \times G_0(r, r_0, \omega_0) \cdot m(\omega_0),$$  \hspace{1cm} (1)$$

and

$$H_0(r, \omega_0) = \{ k_0^2 + \nabla_r \nabla_r \} G_0(r, r_0, \omega_0) \cdot m(\omega_0),$$  \hspace{1cm} (2)$$

where $k_0 = 2\pi/\lambda_0$ is the wave vector in vacuum and $G_0(r, r_0, \omega_0) = \exp(ik_0|r - r_0|)/|r - r_0|$ represents the scalar Green function. From these two equations, we can define two field susceptibilities:

$$E_0(r, \omega_0) = S^{EH}(r, r_0, \omega_0) \cdot m(\omega_0),$$  \hspace{1cm} (3)$$

and

$$H_0(r, \omega_0) = S^{HH}(r, r_0, \omega_0) \cdot m(\omega_0)$$  \hspace{1cm} (4)$$

in which the dyadic tensors $S^{EH}(r, r_0, \omega_0)$ and $S^{HH}(r, r_0, \omega_0)$ are constructed by identification with equations (1) and (2). For the mixed dyad $S^{EH}(r, r_0, \omega_0)$ this identification yields the expression of the nine analytical components:

$$S^{EH}(r, r_0, \omega_0) = ik_0 \begin{pmatrix}
0 & -\frac{\partial \phi_0}{\partial r_z} & \frac{\partial \phi_0}{\partial r_y} \\
-\frac{\partial \phi_0}{\partial r_y} & 0 & -\frac{\partial \phi_0}{\partial r_z} \\
\frac{\partial \phi_0}{\partial r_z} & \frac{\partial \phi_0}{\partial r_z} & 0
\end{pmatrix}.$$  \hspace{1cm} (5)$$

Equations (1) and (2) define the so-called illumination field. Since the materials considered in this article do not directly respond to the optical magnetic field, the coupling with the nanoparticle is entirely described by the first equation. A complete theoretical investigation of this illumination mode requires the accurate computation of the optical field distribution inside the nanostructure for every location $r_0$ of the magnetic dipole. As discussed in the literature, the recent developments of real space approaches for electromagnetic scattering and light confinement established powerful tools for the calculation of the electromagnetic response of complex mesoscopic systems to arbitrary illumination field. Particularly, the technique of the generalized field propagator described in reference provides a convenient basis to derive the electromagnetic response of an arbitrary system to a great number of different external excitation fields. Our approach is based on the computation of a unique generalized field propagator $K(r', r^', \omega_0)$ that contains the entire response of the nanostructure to any incident electric
field \( \mathbf{E}_0(r''', \omega_0) \). Consequently, the self-consistent electric field \( \mathbf{E}(r_0, r'', \omega) \) created inside the nanosystem by a magnetic dipole located at \( r_0 \) can be written as:

\[
\mathbf{E}(r_0, r'', \omega) = \int_v \mathcal{K}(r', r'', \omega_0) \cdot \mathbf{E}_0(r'', \omega_0) \, dr'',
\]

in which the integral runs over the volume \( v \) of the particle. As demonstrated in reference 34, the dyadic \( \mathcal{K} \) writes

\[
\mathcal{K}(r', r'', \omega_0) = \delta(r' - r'') + \mathcal{S}(r', r'', \omega_0) \cdot \chi(r'', \omega_0),
\]

where \( \delta \) is the three-dimensional Dirac function, \( \mathcal{S}(r', r'', \omega_0) \) is the optical field-susceptibility tensor of the nanostructure of electric susceptibility \( \chi(r'', \omega_0) \).

Equation (6) gives access to the electric field inside the nanostructure and therefore to the polarization \( \mathbf{P}(r_0, r'', \omega_0) = \chi(r'', \omega_0) \cdot \mathbf{E}(r_0, r'', \omega_0) \) induced for each position \( r_0 \) of the magnetic dipole. The magnetic field generated outside of the particle can then be calculated by introducing the second mixed field-susceptibility \( \mathcal{S}^{\mathcal{S} \mathcal{E}}(r', r'', \omega_0) = \mathcal{S}^{\mathcal{E} \mathcal{S}}(r', r'', \omega_0) \):

\[
\mathbf{H}(r_0, r, \omega_0) = \int_v \mathcal{S}^{\mathcal{S} \mathcal{E}}(r', r'', \omega_0) \cdot \mathbf{P}(r_0, r', \omega_0) \, dr',
\]

which, in a concise form, leads to:

\[
\mathbf{H}(r_0, r, \omega_0) = \mathcal{S}^{\mathcal{E} \mathcal{S}}(r', r_0, \omega_0) \cdot \mathbf{m}(\omega_0),
\]

where \( \mathcal{S}^{\mathcal{E} \mathcal{S}}(r, r_0, \omega_0) \) defines the magnetic field susceptibility associated with the nanostructure (\( p \)):

\[
\mathcal{S}^{\mathcal{E} \mathcal{S}}(r, r_0, \omega_0) = \int_v \int_v \mathcal{S}^{\mathcal{S} \mathcal{E}}(r', r_0, \omega_0) \cdot \chi(r', \omega_0) \cdot \mathcal{K}(r', r_0, \omega_0) \cdot \mathcal{S}^{\mathcal{E} \mathcal{S}}(r_0', r_0, \omega_0).
\]

Here, the dot “.” signifies the matrix product. This general relationship, derived from the theory of linear response, brings to light the complex link between the electrical response of matter (contained in \( \chi \) and \( \mathcal{K} \)) and the magnetic response of vacuum, through the mixed propagators \( \mathcal{S}^{\mathcal{E} \mathcal{S}} \) and \( \mathcal{S}^{\mathcal{S} \mathcal{E}} \). The combination of these response functions shows in a concise way how a nanostructure, which originally does not possess any magnetic response in the optical spectrum, can nevertheless yield a magnetic–magnetic response. Equation (10) summarizes with mathematical clarity the back-action of the electromagnetic near-field on a magnetic quantum emitter via the curl of the electric field, mediated by the presence of a non-magnetic nanostructure.

### III. MAGNETIC DIPOLE DECAY-RATE CLOSE TO SMALL DIELECTRIC PARTICLES

Equation (10) allows us to obtain a general expression for the decay rate \( \Gamma_m(r_0, \omega_0) \) associated with a magnetic dipole transition of amplitude \( m_{eg} \):

\[
\Gamma_m(r_0, \omega_0) = \Gamma_m^0(\omega_0) \times \left\{ 1 + \frac{3}{2k_0^2} \mathbf{u} \cdot \text{Im}(\mathcal{S}^{\mathcal{E} \mathcal{S}}(r_0, r_0, \omega_0)) \cdot \mathbf{u} \right\},
\]

where \( \Gamma_m^0(\omega_0) = 4k_0^2 m_{eg}^2 / 3h \) represents the natural decay rate of the magnetic transition and \( \mathbf{u} \) labels the dipole orientation.

The next objective of this article is to supply a full analytical treatment of \( \Gamma_m(r_0, \omega_0) \). To achieve this goal, we deliberately reduce the physical model to a simple two–level system coupled to a single spherical nanoparticle as shown in figure 2-a. We have chosen to illustrate our method with dielectric materials as they offer an interesting alternative to metals with reduced dissipative losses and large resonant enhancement of both electric and magnetic near-fields.

In this case, a set of simple analytical equations can be derived that include all the physical effects mentioned above. Indeed, we have \( \mathcal{K}(r', r'', \omega_0) = \delta(r' - r'') (\mathcal{I} \text{ being the identity tensor}) \), \( \chi(r', \omega_0) = \alpha_e(\omega_0) \delta(r'') \), where \( \alpha_e(\omega_0) \) is the dynamical dipolar polarizability of the sphere, and finally:

\[
\mathcal{S}^{\mathcal{E} \mathcal{S}}(r, r_0, \omega_0) = \alpha_e(\omega_0) \mathcal{S}^{\mathcal{S} \mathcal{E}}(r, 0, \omega_0) \cdot \mathcal{S}^{\mathcal{E} \mathcal{S}}(0, r_0, \omega_0).
\]

We will consider the case \( \omega_0 = 0 \), i.e. \( \mathbf{r} = \mathbf{r}_0 = \omega_0 \), we get the following simple expression when \( \mathbf{r} = \mathbf{r}_0 \) (c.f. equation (11)):

\[
\mathcal{S}^{\mathcal{E} \mathcal{S}}(r, r_0, \omega) = \alpha_e(\omega) A_c k^2 m_{eg}^2 / 3h \frac{k_0^2}{k_0^2 + k^2} \frac{2F_0^2}{F_0^2 + F_1^2} \frac{k_0^2}{k_0^2 + k^2} F_2^2,
\]

where \( F_0 = 2k_0^2 - k^2, F_1 = k_0^2 + k^2, F_2 = k_0^2 + k^2 \).
where \( r_0 = |\mathbf{r}_0| \) and the matrix \( \mathcal{A} \) is defined by:

\[
\mathcal{A} = \begin{pmatrix}
   y_0^2 + z_0^2 & 0 & 0 \\
   0 & z_0^2 & -y_0 z_0 \\
   0 & -y_0 z_0 & y_0^2
\end{pmatrix}.
\]

(14)

In consequence, \( S_{PH}^{HH} \) has the dimension of an inverse volume. A concise expression of the normalized magnetic decay rate \( \Gamma_m = \Gamma_m/\Gamma_m^0 \) can then be deduced by replacing this relation into (11):

\[
\Gamma_m(r_0, \omega_0) = 1 + \alpha_e(\omega_0) \mathbf{u} \cdot \mathcal{A} \cdot \mathbf{u}
\]

\[
\left\{ \sin(2k_0 r_0) \left( -\frac{3k_0}{2r_0^4} + \frac{3}{2k_0 r_0^2} \right) - \cos(2k_0 r_0) \frac{3}{r_0^4} \right\},
\]

(15)

in which the polarization dissipation term \( \text{Im} \alpha_e(\omega_0) \) has been neglected. We set \( x_0 = 0 \) to obtain the most simple equations possible. Adding it as free parameter is straightforward, yet renders the equations (14) and (15) more complex. The case of a single dipolar dielectric sphere presented in figure 2 shows that the contrast patterns are extremely sensitive to the dipole orientation. The contrast is generally positive on top of the particle except when the dipole is aligned perpendicularly to the scanning plane \((x_0, y_0)\) in which case it vanishes, the sphere becoming invisible for the magnetic dipole. Such a peculiar behavior explicitly appears in equations (14) and (15) for small interaction distances, in particular, when the magnetic dipole enters the very subwavelength range corresponding to \( 2k_0 r_0 \ll 1 \). As a second example, we consider in figure 3 a set of \( p \) identical dielectric particles deposited on a transparent substrate positioned at random locations \( \mathbf{r}_i \) \((i = 1 \text{ to } p)\). The optical properties of such a system can be described by first inserting the relation:

\[
\chi(r, \omega_0) = \alpha_e(\omega_0) \sum_{i=1}^p \delta(r - r_i)
\]

(16)

in equation (10) and then in expression (11). The results are presented in figure 3-b and c. When the particles are well-separated from each other, typically by one wavelength or more, they display a similar contrast as the one described in figure 2. This contrast is reinforced when several particles are grouped together. Isolated particles and assemblies of particles are surrounded by pseudoperiodic ripples that reveal the interferences between the emitting magnetic dipole and the sample.

IV. ELECTRIC AND MAGNETIC DIPOLE DECAY-RATE CLOSE TO COMPLEX DIELECTRIC NANO-STRUCTURES

Whereas equations (10) and (11) provide analytical expressions of the decay rate of magnetic dipoles placed close to very simple nano-objects, these equations can be complemented by an adequate discretization of the particle volume to describe light emission from dipoles in the vicinity of nanostuctures of arbitrary geometries. To this end, we numerically implement the complete field susceptibilities \( S^{HE} \) and \( S^{EH} \) in a discretized version of equation (10):

\[
S_{PH}^{HH}({r_0}, \omega_0) = \sum_{i=1}^N V_{cell} \sum_{j=1}^N V_{cell} S^{HE}(r_i, r_j, \omega_0)
\]

\[
\cdot \chi(r_i, \omega_0) \cdot \mathcal{K}(r_i, r_j, \omega_0) \cdot S^{EH}(r_j, {r_0}, \omega_0).
\]

(17)

The sums (indexes \( i \) and \( j \)) run over all \( N \) discretization cells (of volume \( V_{cell} \)) forming the nanostructure. This numerical procedure gives access to the optical response of complex systems, such as the ones described in figure 4. In this example, we have applied this technique to visualize the footprint induced by a perfect square corral composed of 20 dielectric structures in the initially flat ED and MD decay rate maps. The extension of the entire nanostructure is 1.1 \( \mu \)m, the refractive index is \( n_{pad} = 2 \). A modification of the decay rates ranging between 20 and 50 % is obtained when the magnetic dipole is 30 nm above the nanostructures (fig. 4e-d). Although the coupling is more efficient with an electric dipole (fig. 4a-b), especially when it is perpendicular, the coupling of
c) d)

a) b)

e) f)

g) h)

ment, we show in fig. 4e-h a flat silicon structure (TiO$_2$, Si, or even Ge). To demonstrate this enhance-

ture 4), we observe a contrast reversal above the dielectric pads when passing from an electric to a magnetic dipole. 
The striking phenomenon is accompanied by a shift of the fringe pattern inside the corral by half a wavelength. 
Finally, another type of contrast change is observed when the dipole is perpendicular to the sample. In this second 
case, as illustrated by the maps (b) and (d) of figure 4, we move from a highly localized signal around the pads 
(map (b)) to a broader response distributed along the corral rows (map (d)).

\section{V. Evolutionary Optimization of Metal Nano-structures for Maximum Magnetic Decay Rate}

In order to demonstrate the versatility of our model, we couple our numerical framework to an evolutionary optimization (EO) algorithm. EO tries to find optimum solutions to complex problems by mimicking the process of natural selection. Its principal idea is briefly depicted in figure 5a. Our approach to couple EO to numerical simulations is described in more detail in reference 48. For technical information on the implementation and the used algorithm parameters, see the SI. In the supporting information, we also show an additional single- as well as a multi-objective evolutionary optimization problem, based on the decay-rate formalism. In this section, we use the permittivity of gold\cite{ref49} to demonstrate that our formalism is not limited to dielectric materials. The optimization goal is to find a gold nano-structure which maximizes the ratio of magnetic over electric decay rate $\Gamma_m/\Gamma_e$ at a fixed location ($r_0 = (0,0,80) \text{nm}$). This is a particularly tricky scenario, because metals are known to have a far stronger response to electric dipole transitions than to magnetic ones. We use the evolutionary algorithm to optimize the geometry of a planar structure composed of 100 gold pillars (each $20 \times 20 \times 60 \text{nm}^3$), lying on a plane of $1000 \times 1000 \text{nm}^2$ (see figure 5b). We recall here that each subwavelength pillar does not support a direct magnetic response on its own. To render the positioning easier, the possible locations on the plane lie on a discretized grid (steps of 20 nm). The structure is placed in vacuum and the wavelength is fixed at $\lambda = 500 \text{nm}$. We evolve a population of 150 individuals (nano-structures) over 2500 generations. Each of the individuals is a parameter-set consisting of positions for the 100 gold pillars, hence describing one possible structure. We tested the convergence by running the same optimization several times, reproducibly yielding similar structures and values for the decay rate ratio.

The optimum structure found by the EO algorithm is
FIG. 5: (color online) (a) Evolutionary optimization cycle. (b) Sketch of the structure model for optimization: Free parameters are the positions of 100 gold blocks ($B_i$) on the XY plane (in vacuum). (c) Gold structure for optimum $\Gamma_m/\Gamma_e$ contrast at the center ($r_0 = (0, 0, 80)$ nm), found by EO. (d-g) Decay rate analysis of the EO solution. (d) Mapping of the ratio of magnetic and electric decay rate 20 nm above the structure. (e-f) Relative electric and magnetic decay rates above the structure, respectively. (g) Ratio of magnetic and electric decay rate for only the outer part of the structure, also leading to an enhancement of $\Gamma_m$ at the target location. At the bottom of (g) $\Gamma_m/\Gamma_e$ is shown along a profile in the center of the map. (h) Progress of the evolutionary algorithm. A logarithmic colorscale is used for the maps (e-f). All results are computed at $\lambda_0 = 500$ nm and for a dipole orientation along OZ. Scale bars are 200 nm. Mapping (c) is $1000 \times 1000$ nm$^2$, (d-g) are $800 \times 800$ nm$^2$ large (area indicated by a dashed square in c).

shown in figure 5c. Mappings of the decay rate ratio as well as the electric and magnetic decay rates are shown in figure 5d-g. Obviously, the algorithm succeeded in finding a gold nanostructure which significantly promotes magnetic decay at the target position (see Fig. 5d). This is particularly remarkable, because although gold structures easily provide very strong electric dipole decay rate enhancements, the magnetic LDOS is known to be usually very weak in metallic nanoparticles.26

Two effects are being exploited by the optimized structure: The first mechanism is the different confinement of the decay rates for electric and magnetic dipoles close to material. The electric decay rate enhancement in the proximity of the gold pillars is high, but confined to a very small volume around the material. The magnetic decay rate on the other hand is more loosely enhanced around the gold clusters, leading to regions in their vicinity where $\Gamma_e$ is almost not affected, while $\Gamma_m$ still shows significant enhancement (c-f, figures 5e-f). The second effect is a modulation of the decay rate inside a larger resonator due to interference, similar to the corral shown in figures 1 and 4. At $\lambda = 500$ nm, the above presented corral had a maximum of $\Gamma_e$ in its center (see figure 4b and d). In contrast to this, the evolutionary algorithm distributed a fraction of the material (outer, circular structure) such that $\Gamma_m$ is maximum in its center, which can be seen in figure 5g, where the decay rate has been calculated for the isolated outer structure.

We will conclude this section with some considerations on the convergence. One might wonder why the structure does not consist of perfect circles – this would very likely result in even better performance. Concerning this question, we have to keep in mind that $(51 \times 51)!/(51 \times 51 - 100)! = 10^{144}$ possibilities exist to distribute the 100 gold pillars on the available positions on the plane. Yet, the evolutionary algorithm did only evaluate $2500 \times 150 < 4 \times 10^5$ different arrangements. Therefore, the reason why the material is not distributed on perfect circles is the heuristic nature of the evolutionary optimization algorithm. The search for the best structure did simply not converge to the very optimum. Comparing the optimized structure to an idealized version reveals, that the possible improvement in $\Gamma_m/\Gamma_e$ is only in the order of $\approx 1\%$ (see also supporting information). We conclude that, despite the residual disorder in the geometry, the EO algorithm did converge very close to the ideal structure. Hence EO is a promising approach to this kind of problems.

VI. CONCLUSIONS AND PERSPECTIVES

In summary, we have developed a concise theoretical framework to describe the dynamics of light emission from magnetic dipoles located in complex nanostructured environments. This method, based on mixed field-susceptibilities, provides analytical expressions of the decay rate in the case of very simple environments. When
the magnetic dipole is located close to nanostructures of arbitrary geometries, the computation of the MD decay rate involves the discretization of the nanostructure volume, the computation of a generalized propagator and finally the computation the decay rate from mixed field susceptibilities. This versatile framework is well suited to describe the emission of light from emitters involving both electric and magnetic dipole transitions as well as nano-optical processes comprising confined electric and magnetic fields. In addition, our framework is very flexible and can easily be extended. For instance nonlocality effects might be included by following the descriptions of Ref. 50. Thanks to its computational simplicity, the method can also be employed within more complex numerical schemes. We demonstrated this possibility by coupling the magnetic decay rate calculation to an evolutionary optimization algorithm, which we employed to identify the nature of the transition involved in the decay rate in the vicinity of nanostructures. Finally, nanostructures possessing a particularly high contrast regarding dipole orientations could be designed using our evolutionary optimization.

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