Polarization state of the optical near–field

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The polarization state of the optical electromagnetic field lying several nanometers above complex dielectric–air interfaces reveals the intricate light–matter interaction that occurs in the near–field zone. From the experimental point of view, access to this information is not direct and can only be extracted from an analysis of the polarization state of the detected light. These polarization states can be calculated by different numerical methods well–suited to near–field optics. In this paper, we apply two different techniques (Localized Green Function Method and Differential Theory of Gratings) to separate each polarisation component associated with both electric and magnetic optical near–fields produced by nanometer sized objects. A simple dipolar model is used to achieve insight into the physical origin of the near–field polarization state. In a second stage, accurate numerical simulations of field maps complete data produced by analytical models. We conclude this study by demonstrating the role played by the near–field polarization in the formation of the local density of states.

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I. INTRODUCTION

Light interactions with dielectric or metallic surfaces displaying well–defined subwavelength–sized structures (natural or lithographically designed) give rise to unusual optical effects [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12]. The recently observed “light confinement state” in which the light field is trapped by individual surface defects, belongs to this class of phenomena [8]. Although, with usual dielectric materials (silica for example), the local near–field intensity variations observed around the particles (or structures) remains moderate over the optical spectrum (between 10 to 40 per cent of the incident light intensity), these variations can nevertheless be easily mapped with the tip of a Photon Scanning Tunneling Microscope (PSTM) [8]. The images recorded with this technique reveal dramatic changes when passing from the TM (transverse magnetic) to the TE (transverse electric) –polarized modes. In general, TM–polarized light tends to display larger contrast than TE–polarized light. From the experimental point of view, the definition of the polarization direction of the incident and detected intensity must be defined with respect to a unique incident plane. About six years ago Van Hulst and collaborators proposed a clever probe configuration devoted to polarization mapping [13]. These authors performed these measurements by using a combined PSTM/AFM microscope in which detection is implemented by a microfabricated silicon–nitride probe. From this technique, polarization contrast is extracted by changing the polarization directions of both the incident and the detected light. The main findings gathered in this work[13], concern the relative efficiency of the four excitation–detection possibilities (TE/TE,TE/TM,TM/TE, and TM/TM) to record a highly resolved PSTM image. In particular, the efficiency of the TM/TM acquisition mode is well reproduced. Although a complete interpretation of this work requires a realistic numerical implementation of the combined AFM/PSTM probe tip, we can obtain useful information by analyzing the near–field polarization state versus the polarization state of the illumination mode.

In addition, in closely related contexts, the control of the near–field polarization state provides an interesting and versatile tool for generating powerful applications (tunneling time measurements [14], highly–resolved microscopy and spectroscopy [15], surface plasmon resonance spectroscopy of molecular adlayers [16], atom optics [17, 18]). More precisely, in the field of atom optics and interferometry, one is interested in building diffraction gratings that can play the role of beam splitters. Several devices have been successfully realised, ranging from mechanical transmission gratings to light standing waves in free space or evanescent for a prism. For a general review the reader is referred to [16]. To circumvent some theoretical limitations [17] it has been recently proposed [18] to use micrometer sized metallic stripes to shape the evanescent field. A full near-field, metallic/dielectric approach open obviously new perspectives. In particular the spacing period is no longer linked to the atomic optical transition and a reflection structure cannot clug. More, higher harmonics in the optical evanescent field can be tuned to produce for example a blazed atomic
grating. Nevertheless, the optical potential is strongly related to the light field polarisation, which is a further motivation for the present study. We will begin our theoretical analysis with a simple dipolar scheme in which the main experimental parameters (incident angle, optical index, polarization of the incident light, ...) appear explicitly. In a second stage, these results will be completed with an ab-initio approach allowing objects of arbitrary shape to be treated exactly.

II. POLARIZATION OF THE LIGHT ABOVE A SINGLE DIELECTRIC PARTICLE

To illustrate the coupling between a polarized incident wave and a small spherical object lying on the sample, we consider the simple dipolar model depicted in Fig. 1. The substrate modifies the polarizability $\alpha_0(\omega)$ of the particle. We have then

$$\alpha_{\text{eff}}(\mathbf{R}, \omega) = \alpha_0(\omega) \cdot \mathbf{M}(\mathbf{R}, \omega)$$

with

$$\mathbf{M}(\mathbf{R}, \omega) = [1 - \mathbf{S}_s(\mathbf{R}, \omega) \cdot \alpha_0(\omega)]^{-1}$$

where $\mathbf{S}_s(\mathbf{R}, \omega)$ is the nonretarded propagator associated with the bare surface, and $\mathbf{R} = (0, 0, R)$. XOZ is the incident plane.

![FIG. 1: Schematic drawing of the model system used in the present section. A transparent substrate, of optical index $n = c^2$ supports a small dielectric sphere of radius $R$. The system is illuminated in total internal reflection with an incident angle $\theta_0$ and $\mathbf{R} = (0, 0, R)$.](image)

and

$$\alpha_{\text{eff}}(\mathbf{R}, \omega) = \frac{4(n^2 + 1)\alpha_0(\omega)R^3}{4(n^2 + 1)R^3 - \alpha_0(\omega)(n^2 - 1)}$$

where $n$ is the optical index of refraction of the substrate.

A. New field components in the near–field domain

At an observation point $\mathbf{r}$ located above the sample ($z > 0$) and in the immediate proximity of the particle, the incident light field is locally distorted. As illustrated in Ref. 27, these distortions generate not only a profound modification of the intensity level (both electric and magnetic), but also a complete change of the polarization state. At subwavelength distances from the scatterers, we expect therefore to observe the occurrence of new components that were absent in the incident field $\{\mathbf{E}_0(\mathbf{r}, t); \mathbf{B}_0(\mathbf{r}, t)\}$. The physical origin of this polarization transfer can be easily understood if we introduce the two relevant field propagators $\mathbf{S}_0$ and $\mathbf{Q}_0$ that establish the physical link between the oscillating dipole $\mu(t) = \alpha_{\text{eff}}(\mathbf{R}, \omega_0) \cdot \mathbf{E}_0(\mathbf{R}, t)$ and the new near–field state $\{\mathbf{E}(\mathbf{r}, t); \mathbf{B}(\mathbf{r}, t)\}$ generated above the particle:

$$\mathbf{E}(\mathbf{r}, t) = \mathbf{E}_0(\mathbf{r}, t) + \mathbf{S}_0(\mathbf{r}, \mathbf{R}) \cdot \alpha_{\text{eff}}(\mathbf{R}, \omega_0) \cdot \mathbf{E}_0(\mathbf{R}, t)$$

and

$$\mathbf{B}(\mathbf{r}, t) = \mathbf{B}_0(\mathbf{r}, t) + \mathbf{Q}_0(\mathbf{r}, \mathbf{R}, \omega_0) \cdot \alpha_{\text{eff}}(\mathbf{R}, \omega_0) \cdot \mathbf{E}_0(\mathbf{R}, t)$$

where in the near–field zone, i.e. when $|\mathbf{r} - \mathbf{R}| < \lambda_0 = 2\pi c/\omega_0$,

$$\mathbf{S}_0(\mathbf{r}, \mathbf{R}) = \frac{3(\mathbf{r} - \mathbf{R})(\mathbf{r} - \mathbf{R}) - |\mathbf{r} - \mathbf{R}|^2 \mathbf{1}}{|\mathbf{r} - \mathbf{R}|^3}$$

and

$$\mathbf{Q}_0(\mathbf{r}, \mathbf{R}, \omega_0) = \frac{i\omega_0}{c|\mathbf{r} - \mathbf{R}|^3} \begin{pmatrix} 0 & -(z - R) & y \\ z - R & 0 & -x \\ -y & x & 0 \end{pmatrix}$$

The discussion of the two equations (5) and (6) can be made easier when considering specific examples.

B. Illumination with a TE–polarized surface wave

In this incident polarization mode, the electric field is directed along the (OY) axis. We have then $\mathbf{E}_0(\mathbf{R}, t) = (0, E_0(t), 0)$. Let us see what happens when the observation point moves along the diagonal straight line (A–B) schematized in figure 2. For a given $\beta$ angle, the introduction of the position vector $\mathbf{r} = (r \cos(\beta), r \sin(\beta), z)$ (where $r$ varies between $-\infty$ and $\infty$) along the line (A–B), allows the electric field polarization evolution to be
observed when passing over the particle. Each cartesian component can be simply extracted from Eq. (5). This leads to three analytical relations:

\[ E_x(t) = E_0(t) \frac{\alpha_{eff}(R, \omega) 3r^2 \cos(\beta) \sin(\beta)}{|r^2 + (z - R)^2|^{5/2}} \]  

(9)

\[ E_y(t) = E_0(t) \{1 + \alpha_{eff}(R, \omega_0) T(\beta)\}, \]  

(10)

with

\[ T(\beta) = \frac{2r^2 \sin^2(\beta) - r^2 \cos^2(\beta) - (z - R)^2}{|r^2 + (z - R)^2|^{5/2}}, \]  

(11)

and

\[ E_z(t) = E_0(t) \frac{\alpha_{eff}(R, \omega_0) 3r \sin(\beta)(z - R)}{|r^2 + (z - R)^2|^{5/2}}. \]  

(12)

Some interesting features can be deduced from these equations. (i) First, we observe that the polarization is not modified when the observation point is perpendicular the particle center (i.e. when \( r = 0 \)). The set of equations reduces then to

\[ E_x(t) = 0, \]  

(13)

\[ E_y(t) = E_0(t) \{1 - \frac{\alpha_{eff}(0, 0, z, \omega_0)}{|z - R|^3}\}, \]  

(14)

and

\[ E_z(t) = 0. \]  

(15)

Clearly the effective polarizability reduces the electric field magnitude compared to its initial value. This fact explains the observation of contrasted dark zones above small particles lighted with the TE polarization (c.f. Fig. (4)). (ii) Around the particle (when \( R/2 < r < 2R \)) two new components, namely \( E_x(t) \) and \( E_z(t) \), define a new local polarization state. These components vanish again when the observation point moves away from the particle. Similar relations can be derived for the TM–polarized mode from Eq. (5).

To conclude this section, let us examine what happens with the magnetic field part (cf. Eq. 6). Since in the reference system of Fig. (1), the incident magnetic field displays two components different from zero, \( B_0(r, t) = (B_{0x}(t), 0, B_{0z}(t)) \), we can write

\[ B_x(t) = B_{0x}(t) - \frac{i \omega_0(z - R)}{c|r^2 + (z - R)^2|^{3/2}} \alpha_{eff} E_0(t), \]  

(16)

\[ B_y(t) = 0, \]  

(17)

and

\[ B_z(t) = B_{0z}(t) + \frac{i \omega_0 r \cos(\beta)}{c|r^2 + (z - R)^2|^{3/2}} \alpha_{eff} E_0(t). \]  

(18)

Unlike what happens with the electric field, the particle does not produce new magnetic field components in the near-field. In this case, the polarization change corresponds to a different balance in the initial components. It is important to recall that we have used the dipole approximation to describe the particle-field interaction, i.e. the size of the particle is assumed to be small compared to the wavelength of light. In a more realistic calculation, with nanostructures of characteristic dimension \( \approx 100 \) nanometers this result is not rigorously exact. However, we still expect, in TE polarization mode, a negligible particle contribution to \( B_y \) compared to the total magnetic field intensity.

III. AB INITIO STUDY OF THE NEAR–FIELD POLARIZATION STATE

Analytical results presented in the previous section supply qualitative information about the spatial polarization state distribution. In a recent paper, analysis of polarization effects was proposed in the context of near–field optics in which a limited number of single particles were investigated[26]. Since in many practical situations experimentalists are interested in lithographically designed structures, these preliminary analyses must be completed by \textit{ab initio} procedures for solving Maxwell’s equations.

A. Localized objects

Recently, theoretical modelling in the vicinity of localized objects was performed in the framework of the
field susceptibility method\textsuperscript{[27, 28]}. Today, this method is one of the most versatile and reliable numerical techniques to solve the full set of Maxwell equations for the typical parameters of near-field optics. It works well even for metallic nanostructures (see for example references\textsuperscript{[6, 8, 10]}). This approach (called the Direct Space Integral Equation Method (DSIEM)) is based on the knowledge of the retarded dyadic tensor $S(r, r', \omega)$ associated with a reference system which, in our problem, is a flat silica surface\textsuperscript{[29, 30]}. The numerical procedure considers any object deposited on the surface as a localized perturbation which is discretized in direct space over a predefined volume mesh of $N$ points $\{R_i\}$. In a first step, the electric field distribution $E(R_i, \omega)$ is determined self-consistently inside the perturbations (i.e., the source field). At this stage, a renormalization procedure associated to the depolarization effect is applied to take care of the self-interaction of each discretization cell. The final step relies on the Huygens–Fresnel principle to compute the electromagnetic field $E(r, \omega)$ on the basis of the knowledge of the field inside the localized perturbations $E(R_i, \omega)$. The two main computational steps can be summarized as follows:

(i) Local field computation inside the source field

$$E(R_i, \omega) = \sum_j K(R_i, R_j, \omega) \cdot E_0(R_j, \omega), \quad (19)$$

where $K$ labels the generalized field propagator of the entire system (localized object plus bare silica surface). In the $\{R_i; R_j\}$ representation it is given by

$$K(R_i, R_j, \omega) = \delta_{ij} + v_j S(R_i, R_j, \omega) \cdot \chi(R_j, \omega), \quad (20)$$

where $\chi$ represents the electric susceptibility of the localized object, $v_j$ is the volume of the elementary discretization cell, and $S$ is the field–susceptibility of the entire system. This last quantity is usually computed by solving Dyson’s equation:

$$S(R_i, R_j, \omega) = S(R_i, R_j, \omega) + \sum_k v_k S(R_i, R_k, \omega) \cdot \chi(R_k, \omega) \cdot S(R_k, R_j, \omega), \quad (21)$$

(ii) Electric and magnetic near–field mapping computation around the source field region

$$E(r, \omega) = E_0(r, \omega) + \quad \sum_i v_i S(r, R_i, \omega) \cdot \chi(R_i, \omega) \cdot E(R_i, \omega). \quad (22)$$

and

$$B(r, \omega) = B_0(r, \omega) + \quad \sum_i v_i Q(r, R_i, \omega) \cdot \chi(R_i, \omega) \cdot E(R_i, \omega). \quad (23)$$

In the numerical work to be discussed in this section, the retarded propagators $S$ and $Q$ have been chosen in reference\textsuperscript{[31]}.

The test–object we consider in this section is the word OPTICS engraved at the surface of a TiO$_2$ layer deposited on a silica surface. Intentionally we have chosen a planar structure devoid of any symmetry. In order to gain more insight in the polarization changes occurring around such complex lithographically designed nanostructures, we analyze in Figs. (4)-(7) the electric and magnetic near–field intensities generated by each cartesian component ($E_x^2$, $E_y^2$, $E_z^2$) and ($B_x^2$, $B_y^2$, $B_z^2$). For comparison, the square modulii are also provided.

1. Dielectric materials

The lateral dimensions of the object are given in Fig. 3. The thickness and the optical index of the TiO$_2$ pattern is 20 nm and 2.1, respectively. The wavelength of the incident laser is 633 nm. All fields are computed 10 nm above the surface of the structure, i.e. 30 nm above the glass-air interface. The incident light is a TM/TE–polarized evanescent surface wave traveling along the Ox axis. This illumination condition is used in the Photon Scanning Tunneling Microscope (PSTM). Some general comments can be made about these results. First, all components of both the electric and magnetic fields have been excited in the near-zone. The occurrence of these new components is a pure near–field effect because it is always localized around the structures. In Figs. 3 and 4 we display the electric field part. As predicted in sec-
FIG. 4: Grey scale representation of the electric field distribution computed above the topographical object depicted in Fig. 3. The calculation is performed in the TE-polarized mode and the arrow indicates the propagation direction of the surface wave. Extreme values of the components of the electric field (normalized by the incident field) are: 0.000 (min) and 0.153E-1 (max) for $E^2_x$, 0.674 and 1.67 for $E^2_y$, 0.000 and 0.747E-1 for $E^2_z$, 0.681 and 1.68 for $E^2$. 

FIG. 5: Same as figure 4, but in TM-polarized mode. Extreme values are: 0.206 (min) and 0.436 (max) for $E^2_x$, 0.700E-9 and 0.216E-1 for $E^2_y$, 0.613 and 1.06 for $E^2_z$, 0.908 and 1.31 for $E^2$.

In section 2, we recover the appearance of two additional components, $E_z$ and $E_y$, when the object is excited by a TE-polarized surface wave. In agreement with the PSTM results, numerous regions appear with a dark contrast and a moderate intensity level.

As expected, the excellent image–object relation currently observed in the TM-polarized mode is mainly provided by the field component $E_z$ normal to the object. The two other contributions tend to slightly degrade the total pattern $E^2$ composed by the superposition of the three maps $E^2_x$, $E^2_y$, and $E^2_z$.

The magnetic near-field intensity maps (cf. Figs. 6, 7) also show a significant confinement of the magnetic field over the particle which reverses the contrast with respect to the electric map. Similarly to what happens with the electric field, the role played by the additional components can degrade the topographic information contained in the complete field maps. Notice in Fig. (6) that, as mentioned in section II, the new y-component of the magnetic field is very small compared to the total magnetic field.

2. Metallic materials

In the above formalism, the only parameter distinguishing metallic from dielectric objects is the linear susceptibility $\chi(r', \omega)$. Alternative procedures can be adopted to describe the metallic susceptibility. For example, a direct route would consist in expanding the susceptibility in a multipolar series around the geometrical
FIG. 7: Same as figure 6, but in TM–polarized mode. Extreme values are 0.109E-7 (min) and 0.856E-2 (max) for $B_x^2$, 0.710 and 1.11 for $B_y^2$, 0.107E-7 and 0.190E-1 for $B_z^2$, 0.711 and 1.11 for $B^2$.

center of the metallic particle. While this scheme allows non–local and quantum size effects to be included, it is nevertheless restricted to simple particle shapes (spheres, ellipsoids, etc.). When dealing with spherical metallic clusters having a typical radius below 15 nm this description is mandatory and can be easily included in the DSIEM formalism[32]. For the applications discussed in this paper, involving lithographically designed metallic structures larger than this critical size, we can adopt the discretization of $\chi(r', \omega)$ over all the volume occupied by the particle. In this case, the local susceptibility is just related to the metal optical index $n$ by the relation[9]:

$$\chi(r', \omega) = \frac{(n^2(\omega) - 1)}{4\pi}$$

(24)

In the visible range, the numerical data for describing both real and imaginary parts of $n$ have been tabulated by Palik[33] for different metals. We present in Fig. 8 a gray scale representation of the electric field distribution computed above the topographical object depicted in Fig. 3. In this application the supported structure is metallic (gold). The incident wavelength is 760 nm. The calculation is performed in the plane $z_0 = 30$ nm in the TM–polarized mode and the arrow indicates the propagation direction of the surface wave. Extremal values of the components of the normalized electric field are 0.118E-2 (min) and 3.13 (max) for $E_x^2$, 0.146E-5 and 1.72 for $E_y^2$, 0.237 and 10.1 for $E_z^2$, 0.851 and 10.6 for $E^2$.

longer applicable. But any figure can be decomposed in points (direct space) or Fourier components (reciprocal space). We thus can use two methods whether the real object contain few points or few Fourier components. In the case of periodic gratings, the field distribution can be investigated with this second class of methods [34, 35, 36].

The so–called differential theory of gratings (DTG) was originally developed twenty years ago to predict the efficiencies of one– and two–dimensional diffraction gratings. Based on a rigorous treatment of Maxwell’s equations, this method can also be used efficiently to determine the optical near–field scattered by three dimensional periodic objects. In the following subsection, in order to avoid a complete presentation of this well–established technique, we will only summarize the essential steps of the computational procedure.

As in previous sections, we are interested in the electromagnetic near-field diffracted above objects engraved on an interface illuminated by total internal reflection. When using the DTG method[34], the electromagnetic field above the grating can be expanded in a Fourier series

$$A(r) = \sum_{p=-\infty}^{+\infty} \sum_{q=-\infty}^{+\infty} A_{p,q} e^{i(p\alpha + q\beta)k_{p,q} \cdot r}$$

(25)

where $r = (l, z) = (x, y, z)$. $A(r)$ represents either the
The set of wave vectors $k_{p,q} = (k_{p,q})$, associated with the harmonic $(p,q)$ obey the well-known dispersion equation
\[ k_{p,q}^2 + \gamma_{p,q}^2 = n^2 k_0^2, \tag{26} \]

where $k_0$ denotes the incident wavelength, $n$ the refractive index of the medium, and $\gamma_{p,q}$ the effective propagation constant. The coefficients $\gamma_{p,q}$ are determined by the boundary conditions at the surface of the sample and are obtained by solving the transcendental equations (26). The physical meaning of $\gamma_{p,q}$ is that it corresponds to the real part of the propagation constant of the scattered field, while the imaginary part is associated with the absorption of the incident wave.

In a general way, the six components of the electromagnetic field $\mathbf{A}(r)$ can be deduced from two independent parameters usually named the principal components. Let us choose, for example, the $y$-components $E_y(r)$ to determine the principal components. It is a simple matter to show that the Fourier $y$-components of the field just above the surface of objects can be expressed as a linear combination of the $y$-components of the incident field:
\[ \begin{align*}
E_{y,p,q} &= T_{EE}^{pq} E_{0y} + T_{EB}^{pq} B_{0y}, \\
B_{y,p,q} &= T_{BE}^{pq} E_{0y} + T_{BB}^{pq} B_{0y}.
\end{align*} \tag{28} \]

The transmission coefficients $T_{EE}^{pq}$, $T_{EB}^{pq}$, $T_{BE}^{pq}$ and $T_{BB}^{pq}$ describe the coupling between the electric and magnetic harmonics composing the scattered and the incident field. These coefficients depend both on the geometry of the sample and on the angular conditions of incidence but not on the polarization of the incident light. The polarization of the incident plane wave is controlled by the values of $B_{0y}$ and $E_{0y}$. From a numerical point of view, the transmission coefficients are obtained by the inversion of a complex square matrix whose dimension is $2N_T \times 2N_T$ (where $N_T$ is the total number of harmonics used to describe the scattered field in Eq. (23)). Columns of this matrix contain the Fourier $y$-components of the electromagnetic field which would have illuminated the periodical objects in order to obtain a pure harmonic field $(p,q)$ just above the nanostructure. A detailed description of the calculation of the matrix elements can be found in Refs. [44, 45]. With Eq. (28), we can calculate all the Fourier components of electric and magnetic fields just above the objects, which are used as initial conditions to obtain the field anywhere. General remarks about contrast, relative intensities and image-object relation have been made previously (see sections IIIA and IIIB), therefore we only highlight the specific electromagnetic properties of periodic structures.

We studied a lattice of $100 \times 100 \times 100 \text{ nm}^3$ $TiO_2$ dots, separated by 150 nanometers. A strong localization of the electric field appears above the pads in TM-polarized mode or between the pads in TE-polarized mode. More-
over, a careful analysis of the different components shows that it is possible to create a particular field map such as field lines oriented along y-axis \( (E_y \text{ in TE-mode}) \) or x-axis \( (E_x \text{ in TM-mode}) \), or periodic field spots with very different characteristics (spot size, periodicity and shape) considering the other components in the two polarization modes. These particular field components distribution could have a great interest for the interaction of cold atoms with optical evanescent waves \[37, 38\].

**IV. LOCAL DENSITY OF STATE AND POLARIZATION EFFECTS**

Unlike what happens with electronic surface states, the Local Density of Photonic States (the so-called photonic LDOS) contains information related to the polarization of the excitation field. It is well established that the density of states near surfaces plays a significant role in near-field physics \[12\]. In particular, the photonic LDOS is a useful concept for the description of fluorescence decay rate in the very near-field \[12\] and could help in understanding image formation produced by illuminating-probe SNOM \[40\]. By referring to the electric field, we deduce the optical LDOS from the field-susceptibility of the entire system (plane surface plus supported nanostructures, see section II A) \[40, 41\]

\[
\rho(r, \omega) = \frac{1}{2\pi^2 \omega} Im [Tr S(r, r, \omega)] .
\] (29)

In this expression, the optical LDOS is related to the square modulus of the electric field associated with all electromagnetic eigenmodes of angular frequency \( \omega \). Because of the vectorial character of electromagnetic fields, it is very useful to introduce three polarized optical LDOS also called partial LDOS, so that \[40\]

\[
\rho_i(r, \omega) = \rho_x(r, \omega) + \rho_y(r, \omega) + \rho_z(r, \omega), \quad i = x, y, z \tag{30} 
\]

\[
\rho_i(r, \omega) = \frac{1}{2\pi^2 \omega} Im S_{ii}(r, r, \omega), \quad i = x, y, z \tag{31} 
\]

The three different polarized optical LDOS computed over a pattern made of three dielectric cylinders of optical index 2.1 are represented on figure 14. At this stage, we can make some general remark. In the configuration investigated in figure 14 we can verify that both X–LDOS and Y–LDOS are almost identical with just a rotation of \( \pi/2 \) between them. Nevertheless, this properties does not subsist any more if we reinforce the optical coupling between the dielectric posts. More precisely, certain arrangements of nanoscale pillars (circular or ellipsoidal) can force light waves into states generated by the collective coupling between the pillars leading to significant difference between X–LDOS and Y–LDOS maps.

Moreover, although optical LDOS characterizes the spectroscopic properties of an electromagnetic system independently of the illumination mode, it is interesting to note the strong analogy between the dark contrast over the pads obtained for Y–LDOS and the dark contrast that appears in the case of s-polarized illumination mode (i.e. incident electric field along the y-axis) observed in section II B. In addition, as in the case of optical near-field maps discussed in the previous section, the polarized LDOS’s can display significant discrepancies relatively to the shapes of the underlying objects.

Finally, let us note that partial LDOS’s are not only a powerful mathematical tool but can easily be linked to the physical properties of electromagnetic systems. The most famous example is the fluorescence lifetime \( \tau \) of a molecule near an interface which critically depends on the spatial LDOS variation \[42, 43, 44\].

**V. CONCLUSION**

On the basis of both simple analytical model and sophisticated 3D Maxwell’s equations solvers this paper has focussed on the unusual behaviour of the light polarization in the near-field. When subwavelength patterned objects are excited by a surface wave of well-defined polarization, a complex rotation of the light polarization state can be expected in the near zone. This phenomenon localized around the scatterer is a typical near-field effect. The occurrence of new components is more pronounced in the electric field than in the magnetic part. Subwavelength features are present in all components but with very different energy levels.

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