



# All-Dielectric Nanoresonators for $\chi^{(2)}$ Nonlinear Optics

Carlo Gigli<sup>1</sup>, Giuseppe Marino<sup>1</sup>, Adrien Borne<sup>1</sup>, Philippe Lalanne<sup>2</sup> and Giuseppe Leo<sup>1\*</sup>

<sup>1</sup> MPQ, Université de Paris, CNRS, Paris, France, <sup>2</sup> LP2N, Institut d'Optique Graduate School, CNRS, Univ. Bordeaux, Talence, France

Metal-less nanophotonics offers new opportunities for non-linear optics with respect to optical waveguides and microresonators, taking advantage of the progress within nanofabrication to boost the development of subwavelength Mie-type structures. Here, we review recent results on second harmonic generation with semiconductor nanoresonators, focusing on their scattering features in terms of efficiency and control over radiation patterns. First, two theoretical models are comparatively discussed with a view to possible improvements in analysis and design. Then, some relevant experiments are reported, and the origin of the  $\chi^{(2)}$  generation is discussed, outlining the main open topics to investigate in the near future and the advantages offered by these nanostructures to the development of novel photonic devices.

## OPEN ACCESS

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### \*Correspondence:

Giuseppe Leo  
giuseppe.leo@univ-paris-diderot.fr

### Specialty section:

This article was submitted to  
Optics and Photonics,  
a section of the journal  
Frontiers in Physics

Received: 19 June 2019

Accepted: 29 November 2019

Published: 20 December 2019

### Citation:

Gigli C, Marino G, Borne A, Lalanne P  
and Leo G (2019) All-Dielectric  
Nanoresonators for  $\chi^{(2)}$  Nonlinear  
Optics. *Front. Phys.* 7:221.  
doi: 10.3389/fphy.2019.00221

**Keywords:** nanophotonics, second harmonic generation, Mie resonances, all-dielectric resonators, AlGaAs nanoresonators

## INTRODUCTION

Light manipulation at the nanoscale has been a central topic in nanophotonics for the last two decades. The control of light-matter interaction through subwavelength-resonating elements has led to the possibility of engineering the optical properties of nanopatterned materials, and, in turn, this has led to peculiar phenomena and opportunities for striking applications, including optical magnetism, negative refraction [1, 2], epsilon-near-zero [3, 4], hyperbolic metamaterials [5], sub-wavelength focusing [6], cloaking, wavefront shaping, and flat lenses [7, 8].

Many of these applications originated from the observation of a magnetic response produced by subwavelength structures. In metallic nanostructures, the response to external electromagnetic excitation arises from surface plasmon polaritons or localized surface plasmons. The resonant behavior of these nanostructures allows for the imprint of an abrupt phase discontinuity to propagating light. Control over optical properties allows for the design of 2D arrays of resonators with a spatially varying phase response and, consequently, the fabrication of metasurfaces for beam shaping [7–10].

Nanophotonics has also attracted the interest of the scientific community for intensity-dependent phenomena. Tight field confinement boosts nonlinear generation efficiency in the medium, allowing for the presence of subwavelength nonlinear light sources with amplitude and phase control [11–14]. However, due to their nature, surface plasmons are always accompanied by losses, which limits their application at optical frequencies.

An alternative way to solve this problem is offered by all-dielectric nanostructures. According to Mie theory, high-refractive-index dielectric nanoparticles exhibit strong electric and magnetic resonances, demonstrated for the first time in the visible spectral range with silicon nanoparticles [15, 16]. Control over the phase of the diffracted beam through a 2D array of all-dielectric elements

was already proven in 1998 with local effective index modulation in blazed-binary diffractive elements [17, 18].

The linear resonant behavior of high-refractive-index dielectric nanoparticles is typically obtained via multipolar decomposition of the scattered electromagnetic fields, which consists of the approximation of complex charge distributions with point multipoles [19]. This approach reproduces the observable scattering cross section of the nanoparticles fairly well as it considers the dominant excited multipole moments. However, more accurate methods exist that calculate the scattering cross section from the natural eigenstate basis of the considered open cavity, whose elements are called quasinormal-modes (QNMs) [20–23].

For nonlinear generation processes in the visible range, high-refractive-index nanoparticles are even more advantageous than their plasmonic counterparts due to (a) the absence of ohmic losses and (b) strong field confinement inside the resonator volume, thus enabling the exploitation of the large bulk nonlinearity of these materials. These features are promising for the integration of flat metasurfaces in optoelectronic devices for efficient parametric generation of light with controlled amplitude, phase, and polarization. In this context, the first relevant results have been obtained with third harmonic generation (THG) in silicon nanodisks. In 2014, Shcherbakov et al. demonstrated that local enhancement of an electromagnetic field in silicon nanodisks, due to the magnetic resonance response in the near IR, gives rise to two-order-of-magnitude enhancement of THG with respect to bulk silicon [24], immediately establishing this as an ideal building block for ultra-fast optical switching in integrated photonics [25]. All-dielectric resonators can be thought of as optical cavities with losses. If, on one hand, this makes them attractive as engineerable nonlinear light sources, their nonlinear generation efficiency is, on the other hand, strongly limited by the poor quality factor of resonating modes. In this respect, nanoresonator performances can be boosted by exploiting the multi-mode interference [26], resulting, for example, in Fano resonances [27, 28] or non-radiative anapole modes [29].

Since second-order bulk nonlinear processes are inhibited in centrosymmetric materials like silicon, the quest for high- $\chi^{(2)}$  nonlinear Mie resonators drew the attention on non-centrosymmetric III-V alloys. Among them, promising results have been obtained with aluminum gallium arsenide (AlGaAs) [30, 31] and gallium phosphide (GaP) [32].

This review focuses on three axes of AlGaAs Mie nanoresonators. Firstly, we illustrate two state-of-the-art methods for deriving the linear scattering cross section: multipolar decomposition, and QNM. Secondly, we report on two tools to predict the second harmonic generation (SHG): the first (numerical) consists of finite-elements (FEM) calculations followed by multipolar decomposition, and the second (analytical) relies on Green's functions. Finally, we focus on SHG shaping by highlighting several experimental works.

## SCATTERING PROPERTIES OF ALGAAS MIE-RESONATORS

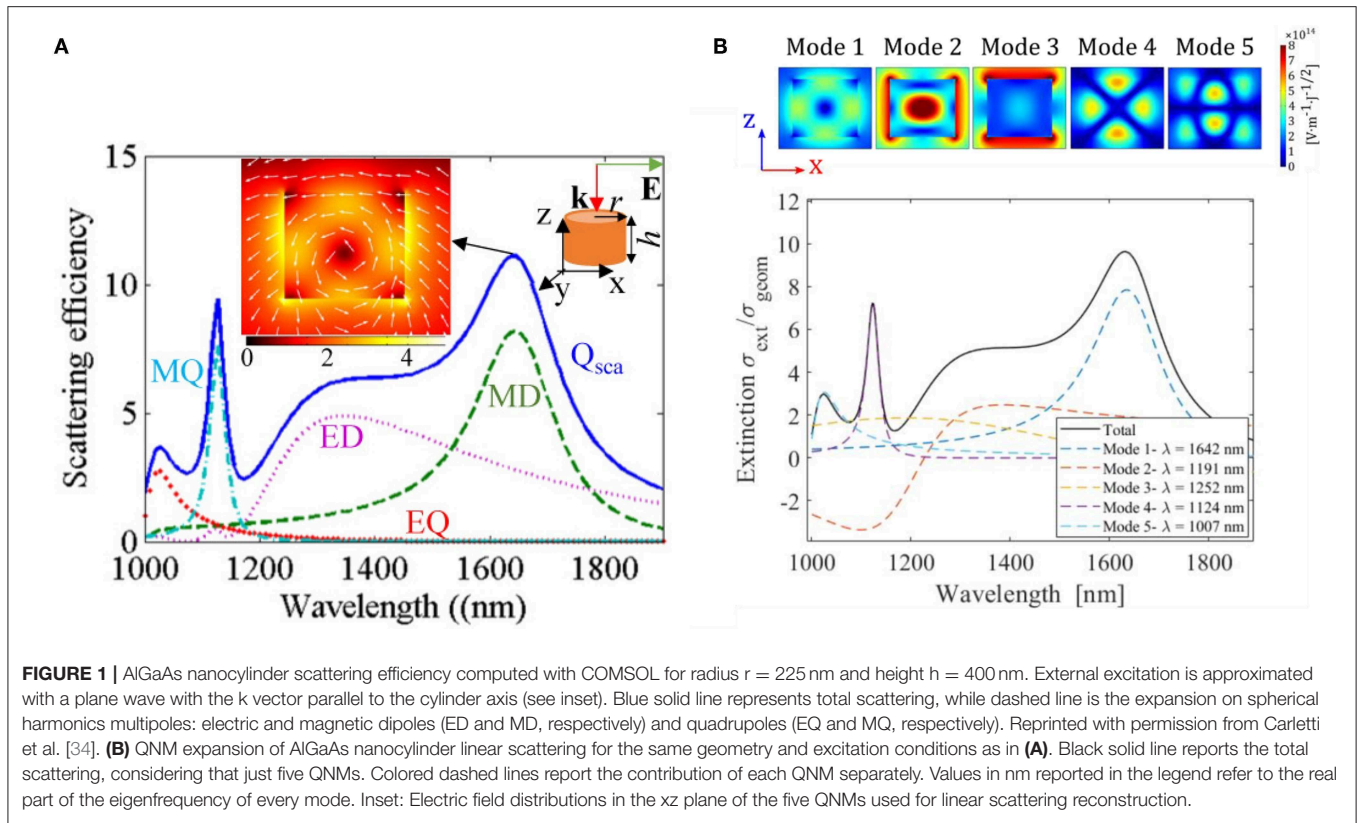
AlGaAs is a zinc-blend alloy exhibiting a high refractive index in the visible/near-IR range ( $n > 3.2$ ), a bandgap that is both larger than GaAs ( $\hbar\omega_g \sim 1.42$  eV), tunable with Al concentration, and has high quadratic nonlinear susceptibility ( $\chi^{(2)} \sim 200$  pm/V) [33]. Its physical properties, combined with a mature technological process, make this material an attractive choice in the framework of III-V nonlinear optical devices. AlGaAs nonlinear Mie-nanoresonators were first investigated by Carletti et al. [34] with  $\text{Al}_{0.18}\text{Ga}_{0.82}\text{As}$  nanocylinders. This alloy composition prevents two-photon absorption in the telecom range while preserving a high refractive index [35]. As an example, take the response of a resonator with radius  $r = 225$  nm and  $h = 400$  nm that is suspended in air to an electromagnetic plane wave with wavevector  $\mathbf{k}$  parallel to the cylinder axis, as is reported in **Figure 1A**.

The inset of the same figure reports the electric field distribution inside the resonator at  $\lambda = 1640$  nm, which testifies to the presence of rotating displacement currents typical of a magnetic dipole. A quantitative understanding of this resonant behavior can be obtained by replacing the complex distribution of charges and currents with an equivalent set of point multipoles generating the same fields [19]. Numerically, this operation is performed by projecting the current distributions originating the scattered field on the basis of spherical harmonics, which leads to the scattering cross section expansion at a given frequency:

$$\sigma_S = \frac{\pi}{k^2} \sum_{l=1}^{\infty} \sum_{m=-l}^l (2l+1) \left[ |a_E(l, m)|^2 + |a_M(l, m)|^2 \right]. \quad (1)$$

The lowest-order multipole contributions to this expansion are reported as dashed lines in **Figure 1A**, which confirms that a strong MD resonance dominates the scattering at  $\lambda = 1640$  nm.

In **Figure 1A**, each resonance of the scattering spectrum is dominated by a multipolar response, which depends on the driving field. At normal incidence, it turns out that the fundamental resonance includes ED and MD contributions. Thus, the multipolar expansion reproduces the scattering cross section of the nanostructure for a given excitation fairly well. However, resonances in the scattering spectrum exhibit contributions not only from the dominant multipole moments that are excited and captured in the decomposition but also from other resonant modes of the considered open cavity. Recently, several numerical tools have been proposed to study radiative and dissipative systems in their proper eigenstate basis [20–23]. In the following section, we demonstrate the deconvolution of extinction spectra of the same system in **Figure 1A** through the method proposed in [23]. In this case, the electromagnetic field in the nanostructure is expanded on the natural basis of the nanoresonator eigenstates, called quasinormal-modes (QNMs),



which are solutions to the source-free Maxwell's equations:

$$\begin{bmatrix} \mathbf{0} & -i\mu_0^{-1}\nabla\times \\ i\epsilon(\tilde{\omega}_m)\nabla\times & \mathbf{0} \end{bmatrix} \begin{bmatrix} \tilde{\mathbf{H}}_m \\ \tilde{\mathbf{E}}_m \end{bmatrix} = \tilde{\omega}_m \begin{bmatrix} \tilde{\mathbf{H}}_m \\ \tilde{\mathbf{E}}_m \end{bmatrix} \quad (2)$$

This is a non-Hermitian eigenvalue problem. QNMs are therefore modes of an open system with a complex eigenfrequency  $\tilde{\omega}_m$ . The scattered electric field can be expanded as [22]:

$$\mathbf{E}_S(\mathbf{r}, \omega) = \sum_m \alpha_m(\omega) \tilde{\mathbf{E}}_m(\mathbf{r}) \quad (3)$$

where  $\tilde{\mathbf{E}}_m(\mathbf{r})$  is the  $m$ -th QNM electric field distribution and the  $\alpha_m$  coefficients are known analytically. The scattering spectrum in **Figure 1A** can be accurately reconstructed using five QNMs, see **Figure 1B**. For the normalization and implementation of this method we refer the reader to [23]. This reconstruction offers several advantages: (a) a robust formalism relying on a proper orthonormal basis; (b) a piece of software implemented in COMSOL Multiphysics; and (c) a gain in terms of computational costs for reconstructing the scattered field irrespective of the incident field.

The proposed case immediately shows the potential of highlighting constructive or destructive interference among QNMs, thereby enabling a good understanding of the physics of the resonating behavior of nanoantennas. It is not surprising to see negative contributions to the total scattering as this is

the footprint of interference between quasinormal modes. It is important to appreciate the change in paradigm from multipole expansion to QNM: in the case of the former, once computed, the scattered field was projected on multipoles distributions; in the latter case, instead, eigenmodes of the structures, i.e., independent of the external source, are analyzed, and then the response to an incident field is retrieved. Furthermore, the electric field distribution of every QNM in the near and far field can be studied separately, which offers a powerful opportunity to apply coupled-mode theory and study mode overlap in the frequency generation processes.

## THEORETICAL ANALYSIS OF SHG IN ALUMINUM GALLIUM ARSENIDE MIE-RESONATORS

The exhaustive comprehension of linear properties enabled the development of models for nonlinear optical phenomena in nanoantennas. In the following section, we have focused on second harmonic generation in AlGaAs resonators. They can be studied numerically with FEM, reducing the analysis to a two-step scattering problem: (1) an external excitation at the fundamental frequency (FF)  $\omega$  is imposed, and electric fields in the nanostructure are retrieved; and (2) a second-harmonic (SH) polarization vector is computed as  $\mathbf{P}^{SH}(\mathbf{r}, 2\omega) = \chi^{(2)}:\mathbf{E}(\mathbf{r}, \omega) \otimes \mathbf{E}(\mathbf{r}, \omega)$ , and Maxwell's equations are solved in the presence of a known current-source distribution

$\mathbf{J}^{SH}(\mathbf{r}, 2\omega) = i2\omega\mathbf{P}^{SH}(\mathbf{r}, 2\omega)$ . At variance with bulk crystals or microresonators, sub-wavelength resonators cannot benefit from phase-matching conditions. Then SHG efficiency depends on electric-field confinement at FF and SH as well as the overlap of these fields mediated by the nonlinear susceptibility tensor [34].

The above requirements imply that having highly resonant modes at FF and SH is necessary, but this is not enough if their distributions do not spatially overlap with respect to symmetry conditions imposed by the  $\chi^{(2)}$  tensor. Indeed, it is immediately possible to highlight that, when moving to shorter wavelengths, the number of excited modes increases. An example is summarized in **Figure 1S**: for [100] grown AlGaAs pillars, crystal symmetry imposes that the magnetic dipole excited at FF by a normally incident beam effectively couples just with SH modes with zero emission in the normal direction. While this constraint might appear as an intrinsic limitation of AlGaAs for applications, it actually illustrates the possibility of exploiting the richness of resonant modes to shape the SHG radiation pattern, polarization, or phase of nonlinear light sources. Taking the same example of a cylindrical resonator with  $r = 225 \text{ nm}$  and  $h = 400 \text{ nm}$ , it was shown that different resonances at SH could be excited by changing the pump polarization or the incident angle [36].

A different strategy to study the SHG efficiency from high-index dielectric nanoparticles (resumed in **Figure 2S**) relies on the Green's function method [37]. This technique, where the SH far field can be calculated via the dyadic Green's function  $\hat{\mathbf{G}}$  of a sphere, has the advantage of showing the selection rules associated with mode coupling based on the symmetry of multipole moments and the crystalline structure. However, it is analytical only in the case of spherical particles and bulk sources. Finally, as an alternative to Green's function, the multipolar coefficients of the SH fields can also be calculated using the Lorentz lemma [38].

## SHAPING SHG FROM ALGAAS NANOANTENNAS: MODEL AND EXPERIMENTAL OBSERVATIONS

The numerical model treated in the previous section enables the investigation of resonance overlap at FF and SH to optimize the nanoantennas design for enhanced SHG. Although the theoretical analyses reported above refer to air-suspended nanocylinders, two slightly different types of (100) high-contrast AlGaAs nanoantennas on a low refractive index have been studied the most: (a) a monolithic device on an aluminum oxide substrate with  $n \approx 1.6$  [30] and (b) a device embedded in BCB with  $n \approx 1.35$  [31]. In particular, it was shown [30] that by tuning a magnetic dipole resonance at FF in the telecom range (around  $\lambda_{FF} = 1.55 \mu\text{m}$ ), a SHG efficiency up to  $6 \cdot 10^{-5}$  could be expected for a nanocylinder pumped with a field intensity of  $1 \text{ GW}/\text{cm}^2$  at normal incidence. This numerical prediction was confirmed experimentally. Please notice that no SHG occurs under normal incidence on (100) bulk zinc-blend crystals, other than a contribution from surface  $\chi^{(2)}$  that is negligible for FF below half bandgap. Furthermore, experimental results for

nanocylinders with radii ranging from 175 to 225 nm pumped at telecom wavelengths displayed good agreement with a theoretical model considering just bulk  $\chi^{(2)}$  contribution. A summary of those results is reported in **Figure 3S**, and the experimental setup is sketched and detailed in **Figure 4S**. This outcome is all the more interesting in that it highlights one of the advantages of dielectric structures over their plasmonic counterparts: strong field confinement inside the volume of the resonator allows for the exploitation of the large bulk nonlinear susceptibility as it yields to about three-order-of-magnitude SHG enhancement with respect to metallic antennas reported in [39] ( $\gamma_{SHG} \cong 1.5 \times 10^{-7} \text{ W}^{-1}$  for the former, instead of  $\gamma_{SHG} \cong 5 \times 10^{-10} \text{ W}^{-1}$  for the latter).

The same outcome also holds for polarization properties, as summarized in **Figure 5SA** where the multipolar expansion based on FEM calculations shows that, as the cylinder radius is varied, SHG is dominated by entirely different resonances. More specifically, as the radius is increased, SH polarization passes from co-polarized to cross-polarized with respect to the polarization of the pump [40].

Similar results regarding the polarization properties of SHG in AlGaAs nanocylinders were detailed by Camacho-Morales et al. [31] (**Figure 5SB**). In that case, the resonators were embedded in a transparent benzocyclobutene (BCB) layer with refractive index  $n \approx 1.35$ , allowing for both a higher refractive index contrast and the possibility of characterizing the device in transmission. In this case, back-focal-plane imaging, enabled via the insertion of a Bertrand lens in the camera detection line, highlighted a strong correlation between the emission direction and SH polarization state. Forward and backward emissions revealed the same features, and, in both cases, the experimental results confirmed full-wave numerical predictions neglecting the contribution of surface  $\chi^{(2)}$ . The comprehension of single element SHG properties, in terms of polarization and directionality, successfully enabled the development of antenna arrays for the controlled generation of SH polarization states [41].

A quite different landscape was explored by Liu et al. [42], which involved working with GaAs resonators on aluminum-gallium oxide. In this case, the dimension of nanoantennas were smaller in order to ensure a magnetic dipole resonance around a pump wavelength of 1020 nm, as detailed in **Figure 6S**.

In contrast to the previous cases, the SH polarization pattern could not be exhaustively explained when considering just bulk  $\chi^{(2)}$  contribution. This is the case particularly when it involves resonances at FF that exhibit strong field intensity close to the resonator surface or when the SH signal is above the energy bandgap, as is displayed in **Figure 6S**, and the SHG contribution from the bulk is strongly absorbed. On the other hand, the introduction of a surface  $\chi^{(2)}$  contribution remains quantitatively challenging due to the unexplored value of this quantity, thereby urging for further and deeper investigations.

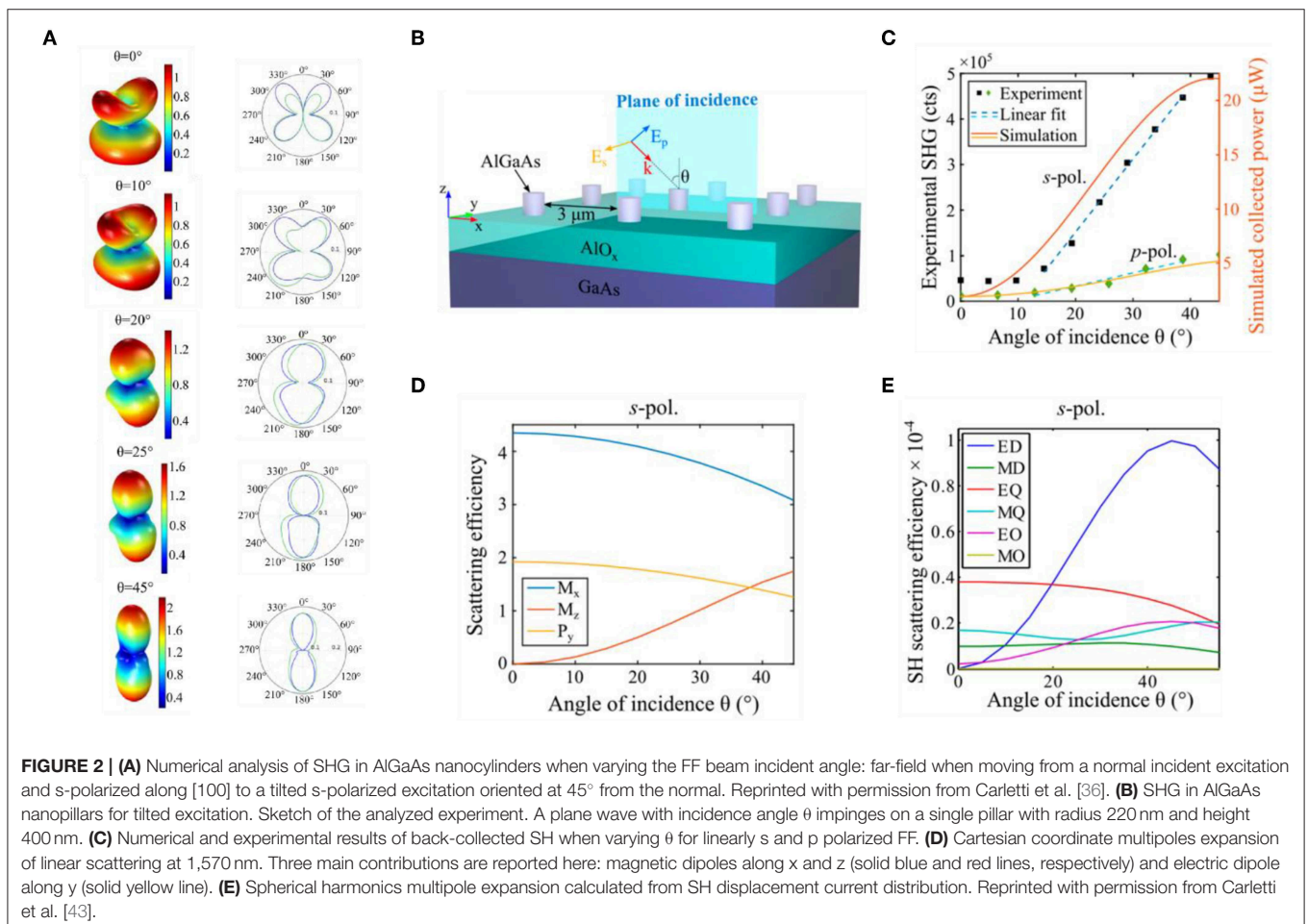
Even if these reviewed works confirm three-order-of-magnitude higher SHG efficiency in AlGaAs nanocylinders compared to plasmonic structures [39], SH normal emission is forbidden due to the symmetry of the nonlinear tensor and the nanoparticle shape. This might compromise the development of integrated optical devices. As shown in previous sections, several

strategies exist to overcome the limitations of  $\chi^{(2)}$  symmetry in [100] grown AlGaAs.

In reference [36], it was numerically predicted that a tilted excitation breaks the electric field symmetry inside the resonator, thereby redirecting SH emission. In that case, for a tilted pump wavevector, different resonating modes exhibiting, for example, normal emission dominate the scattering. This highlights one of the possible mechanisms used to control the SH emission pattern of the nanoscatterer by playing with the excitation of different resonating modes or on the interference between them. These numerical investigations proved that there are a large number of features offered by subwavelength dielectric particles, which rely on the high density of leaky resonant modes in this region of the spectrum. This idea was adopted in Carletti et al. [43] to study the control of the emission pattern of monolithic AlGaAs nanocylinders on an AlOx substrate in the backward direction (see **Figure 2A**). It was shown that changing the incident angle had a strong impact both on multipoles dominating the FF and on SH resonator scattering properties. In particular, with an experimental setup similar to that one of **Figure 4S**, an s-polarized beam at 1570 nm with normal incidence was used to excite an in-plane MD resonance of a 220 nm radius and 400 nm height AlGaAs nanocylinder. When moving to higher

angles thanks to the off-axis lateral translation of the microscope objective, the out-of-plane MD contribution increases, breaking the field symmetry in the resonator. At the SH frequency, this results in a change from a mainly “electric-quadrupole like” field distribution with radiation lobes at large angles and a minimum along the normal axis to an “electric-dipole like” one with two main lobes in the forward and backward directions that point along the cylinder axis (see **Figure 2A**). Exploiting these features, the backward-collected SH can be increased by about one order of magnitude with respect to the two-lobe SH field generated upon normal incidence of the FF pump. Conversely, a p-polarized beam would excite, in nearly the same way, the different FF resonances regardless of the incident angle that was chosen. This results in a quasi-invariant emission, meaning a small enhancement in collected power (see **Figures 2B–E**).

A different approach that maintains normal excitation consists of the integration of a dielectric grating around a nanoparticle so as to control the phase of the nonlinear generation [44]. As explained before, SHG in large-angle lobes is detrimental for the use of on-axis applications. Instead of working on nanoantenna resonances, it is possible to make the in-plane nonlinear emission interact locally with a diffraction grating to reshape the far field (**Figure 7S** for sample images



and experimental results). Due to the symmetry of nonlinear displacement currents, the resonator emission at SH exhibits two main lobes with a  $\pi$ -phase shift, meaning that simply redirecting them to the normal would result in destructive interference. The problem is solved by introducing an asymmetry in grating design in order to further  $\pi$ -shift one lobe with respect to the other. Exploiting this technique an increase by two order of magnitude in the backward collected power with respect to the bare nanocylinder was demonstrated.

An alternative solution to the absence of normal SHG is to rotate the AlGaAs crystalline structure, and thus the  $\chi^{(2)}$  tensor, by fabricating nanoantennas from (111)-GaAs [45]. This change of symmetry of the crystalline structure allows superior forward directionality compared to its (100)-counterparts. The shaping of the SHG radiation pattern via FF polarization was also shown in this case (results are summarized in **Figure 8S**).

## CONCLUSION

Advances in integrated photonics require nonlinear compact devices, like metasurfaces, with high conversion efficiencies and tailorable emission features. After having drawn the attention of the scientific community on the subject of linear applications, nanopatterned dielectric materials have become promising also for nonlinear optics. Here, we reviewed recent progress in high-index dielectric  $\chi^{(2)}$  nanoresonators, which proved advantageous over their plasmonic counterparts because of lower optical losses and a higher conversion efficiency stemming from the overlap of cavity modes with a strong bulk  $\chi^{(2)}$ . Since their multi-resonant behavior enabled the background-free nonlinear displacement current distribution to be tailored locally, a complete understanding of their features is imperative for the SH wavefront design. In this respect, we benchmarked the numerical calculations of the linear scattering cross section against the QNM theory. The latter provides complementary physical insight into Mie-like

resonances in complex structures that cannot be studied analytically. We then focused on fully vectorial numerical calculations, which allowed to model nonlinear effects in good agreement with experimental results. In conclusion, we would like to stress that a better understanding of the relative contribution of bulk vs. surface nonlinear susceptibility would significantly help future progress regarding subwavelength frequency conversion devices. Finally, when compared to their guided-wave counterparts, nanoresonators offer a larger number of spatial degrees of freedom for the engineering of nonlinear generation processes. The development of more powerful design tools, combined with the technological advances of the last years, will probably lead to the creation of unprecedented nonlinear photonic devices.

## AUTHOR CONTRIBUTIONS

CG and GM wrote the paper draft. GL was the scientific supervisor. All the others have contributed to its evolution to the final form.

## FUNDING

GL and PL acknowledge the NOMOS project (ANR-18CE24-0026) for financial support.

## ACKNOWLEDGMENTS

The authors thank C. De Angelis, L. Carletti, D. Rocco, and T. Wu for fruitful discussions.

## SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fphy.2019.00221/full#supplementary-material>

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