Enhanced Second-Harmonic Generation with Structured Light in AlGaAs Nanoparticles Governed by Magnetic Response


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We employ structured light for the second-harmonic generation from subwavelength AlGaAs nanoparticles that support both electric and magnetic multipolar Mie resonances. The vectorial structure of the pump beam allows addressing selectively Mie-resonant modes and control the strength of the generated nonlinear fields. We observe experimentally the enhancement of the second-harmonic generation for the azimuthally polarized vector beams near the magnetic dipole resonance, and match our observations with the numerical decomposition of the Mie-type multipoles for the fundamental and generated second-harmonic fields.

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1. INTRODUCTION

The recently emerged field of dielectric Mie-resonant meta-optics is driven by the studies of dielectric nanoparticles with a high refractive index, and it shapes new active research directions of nanoscale resonant nanophotonics and dielectric nanoantennas [1, 2]. All-dielectric meta-optics employs subwavelength dielectric Mie-resonant nanoparticles as “meta-atoms” for creating highly efficient optical metasurfaces and metadevices [3]. The term “meta-optics” is usually employed to emphasize the importance of the optically induced magnetic response of subwavelength dielectric structures, which are expected to complement or even replace plasmonic components for potential applications at the nanoscale.

High-index dielectric nanoparticles can support both electric and magnetic Mie-type resonances in the visible and near-infrared spectral ranges, and they can easily be tailored by the nanoparticle geometry. The Mie-resonant silicon nanoparticles have recently received considerable attention for applications in nanophotonics and metamaterials [1] including optical nanoantennas, wavefront-shaping metasurfaces, and nonlinear frequency generation. Importantly, the resonant excitation of strong multipolar Mie-type localized modes may also lead to the enhancement of electric and magnetic fields in such structures and drive novel nonlinear effects.

Second-harmonic (SH) generation is one of the most important nonlinear processes in optics. For this parametric process, the frequency of an incident light beam is doubled inside of a nonlinear crystal. Second-harmonic generation is employed for many applications, including laser sources and nonlinear spectroscopy. Remarkably, harmonic generation processes at the nanoscale are governed by the resonant properties of nanoscale structures and the mode volume being independent on the phase-matching principle.

In the last few years, the role of nonlinear optically induced magnetic response was intensively addressed in nanophotonics. In particular, Shcherbakov et al. [4] and Melik-Gaykazyan et al. [5] employed silicon nanoantennas and demonstrated that the nonlinear response from silicon nanodisks prevails over the harmonics generated from the bulk silicon, and it becomes possible to reach conversion efficiency high enough for the generated visible light to be observed by naked eye. Generation of different localized Mie-resonant modes can reshape completely the physics of nonlinear effects at the nanoscale [6]. Utilizing the Mie resonances in dielectric nanoparticles has recently been recognized as a promising strategy to gain higher efficiency of nonlinear parametric processes at low modal volumes, and achieve novel functionalities originating from optically induced nonlinear magnetism [7]. Generated multipoles are closely related to the vectorial nature of the pump beam, and

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they can be distinguished in the far-field region by their polarization and modal decomposition [7].

Focused and structured vector beams have already been employed for the harmonic generation, and they have been used in nanophotonics as a versatile tool, in comparison with the linearly polarized light, for all-optical characterization. A comprehensive review of microscopic techniques based on nonlinear optical processes in nanoscale structures and capabilities of nonlinear microscopy has been discussed recently by Bautista and Kauranen [8] who demonstrated that the vectorial focusing of linear and circular polarized (conventional) and radial and azimuthal polarized (unconventional) light can spread the range of capabilities for nonlinear microscopy techniques.

Recently, we have studied the third-harmonic generation by structured (radially and azimuthally polarized) light [9] and, by tailoring the vectorial structure of the pumping light, we have demonstrated a control of both strength and polarization of the excited harmonic fields, also addressing selectively different types of multipolar Mie-resonances.

In this Letter, we employ our earlier approach for third-harmonic-generation experiments and study, for the first time to our knowledge, nonlinear SH spectroscopy of individual Mie-resonant AlGaAs nanoparticles excited with structured light. Earlier, such problems have been considered only for plasmonic nanoparticles and nanoparticle oligomers [8–11]. We observe the substantial enhancement of the SH nonlinear signal generated by an azimuthally polarized pump beam. We supplement our observations by numerical mode-decomposition analysis and reveal the contribution of the modes at the double frequencies, as well as confirm that structured light can be employed as an efficient tool for the enhanced harmonic generation and subwavelength mode control at the nanoscale.

2. NUMERICAL APPROACH

We simulate numerically both linear and nonlinear optical response, and eigenmode spectra of a single Al_{0.2}Ga_{0.8}As nanodisk using three-dimensional electromagnetic simulations implemented with the finite-element method in COMSOL Multiphysics. The nanodisk is placed on a slide glass substrate, all material properties including real losses are imported from the tabulated data [12]. The vector structure of a pump beam is simulated using the nonparaxial representation [13]. We modify the standard formulas to include rigorously the effect of a substrate by adding reflected and transmitted plane waves into the nonparaxial expansion. For calculations, we take the beam waist radius equal to 1.5 \( \mu \)m.

We choose geometrical parameters of an AlGaAs resonator to excite a magnetic multipolar resonance at the fundamental wavelength by an azimuthally polarized pump beam in the considered near-infrared spectral range (see Fig. 1a). The corresponding geometry of excitation by a radially polarized pump beam is shown in Fig. 1b. The height and diameter of a disk are equal to 650 and 935 nm, respectively. By decreasing the disk radius one can shift this resonance spectral position to the blue range of the spectrum. Figure 2 shows scattering efficiency spectra of an AlGaAs nanodisk of the specified parameters in the linear regime. One can see that indeed the azimuthal polarization excites magnetic Mie-type multipolar modes, whilst contrary to the azimuthally polarized excitation, there are not spectral features for the disk pumped by a radially polarized cylindrical vector beam.

The nonlinear optical response is estimated within the undepleted pump approximation [16]. The induced second-order nonlinear susceptibility tensor \( \chi^{(2)} \) corresponds to zincblende crystalline structure of AlGaAs, determining the value of induced polarization \( P^{(20)} \) at the SH frequency

\[
P^{(20)} = \epsilon_0 \chi^{(2)} E^{(0)} E^{(0)},
\]

where \( E^{(0)} \) is the total electric field inside the resonator at the fundamental frequency. We assume the value of non-zero \( \chi^{(2)} \) components equal to 290 pm/V [17].

We perform the nonlinear numerical simulations of the generated SH electromagnetic fields, their multipolar decomposition and disk eigenmode analysis
It should be pointed out that the spectral position of eigenmodes does not depend on the type of the external excitation and is defined by the parameters of the sample only. The dominant contribution to the SH enhancement of the particle pumped by azimuthally polarized light is made by the mode at the fundamental frequency. Moreover, the spectral features of the electric octupole (EO) contribution can be interpreted by disk eigenmode analysis at the double pump frequency. We present the nonlinear calculation results for two different values of the radius to confirm the spectral shifts of optical modes and their dependence on resonator sizes.

Both scattering spectra and SH power spectra are associated with the electric and magnetic multipolar contributions calculated in the spherical representation [18]. The conventional multipolar analysis does not take the effect of the substrate reflection into account, which provides an error in absolute values of the total contribution of multipoles of order of several percent. In both linear and nonlinear multipolar calculations, the total power is calculated over the full solid angle.

3. EXPERIMENTAL RESULTS AND DISCUSSION

Nanoparticles are made of a custom-designed wafer consisted of a GaAs substrate, a 75-nm sacrificial layer of Al$_{0.51}$In$_{0.49}$P and a 650-nm layer of Al$_{0.2}$Ga$_{0.8}$As with a crystalline axis [100]. The etch mask is created by the method of electron beam lithography; then, sequential processes of dry etching to form pillars and selective wet-etching of the sacrificial layer in solution of HCl are applied. AlGaAs disks are picked up from the GaAs substrate and dropped down to a slide glass substrate. The final dimensions of a nanopillar are checked by scanning electron microscopy. The nanodisk diameter is found to be 935 nm, in correspondence with the value we used for numerical calculations.

In order to conduct nonlinear spectroscopy measurements, we construct an experimental setup based on an optical parametric amplifier (Hotlight Systems, MIROPA-fs-M, pulse duration of about 300 fs) pumped by a pulsed Yb laser by High Q Laser GmbH with a repetition rate of 21 MHz and operating wavelength of 1043 nm. We implement a silicon-based q-plate metasurface [19] to create the vectorial structure of a pump beam from a linearly polarized Gaussian laser beam. The q-plate is placed in a tele-sopic system of two lenses (Thorlabs AC254-200-C-ML and AC254-050-C-ML with focal distances of 20 and 5 cm, respectively). By flipping linear polarization from horizontal to vertical using an achromatic half-wave plate (Thorlabs AHWP05M-1600) we switch the pump beam type from azimuthal to radial. There is a longpass infrared filter (Thorlabs FELH1300) after the q-plate and before the sample to clear up the spectrum of a pump beam. The polarization type of a pump beam is checked by an analyzer, which is a combination of a Thorlabs AQWP05M-1600 achromatic quarter wave-plate and a Thorlabs WP25M-UB broadband wire-grid polarizer. The pump beam is observed by a Xenics Bobcat-320 near-infrared InGaAs camera with a 150-mm focal distance achromatic doublet (AC508-150-C-ML). A train of
wavelength tunable femtosecond laser pulses is focused from the front side of the sample mounted on a three-dimensional stage to a beam waist size, which is close to a diffraction limit, by a Mitutoyo MPlanApo NIR objective lens, 100× infrared, 0.70 numerical aperture (NA). Therefore, the fabricated nanoparticles are considered as isolated due to the spacing value of 10 \( \mu m \). The SH signal is collected by an Olympus objective lens MPlanFL N (100× visible, 0.90 NA) and detected by a visible cooled CCD camera (Starlight Xpress Ltd, Trius-SX694) with a 150-mm focal distance achromatic doublet (AC508-150-A-ML). The detected signal is filtered out by a set of filters (a colored glass bandpass filter Thorlabs FGS900 and UV fused silica filter with dielectric coating FELH0650). The SH signal is normalized over a spectral function of the setup which includes filter transmittance, laser power, and detector sensitivity spectra. The origin of the SH signal is verified by the direct measurement of its spectrum (by a visible spectrometer Ocean Optics QE Pro) and its power dependence which is in a quadratic manner with a saturation average power level of about 250 \( \mu W \).

Our experimental results on the SH generation spectroscopy of an individual AlGaAs resonator of the specific geometrical parameters with cylindrical vector beams are shown in Fig. 4. As expected, we observe the enhancement of the SH signal collected by an objective lens (0.9 NA) for the case of a particle which is resonant at the fundamental wavelength and pumped by an azimuthally polarized laser light. The additional local peaks of the experimental SH spectrum are associated with the excitation of disk eigenmodes at the SH wavelengths, particularly, the increased contribution of an EO mode which could be understood by taking into account the imperfections of fabricated resonators as well as changes of a focused pump beam waist diameter by tuning the laser wave-
length (while it is considered as a constant in our simulations). We observe the four-time less enhancement of the nonlinear response provided by a non-resonant in the linear regime particle excited by a radially polarized laser beam; the spectral position of this enhancement corresponds to the calculated eigenmode of the disk at the double fundamental frequency. This result provides a potential tool of the use of radially polarized light to probe the resonances at doubled frequencies for such type of nanoresonators.

4. CONCLUSIONS

We have employed cylindrical vector beams to enhance the SH generation response of a single AlGaAs nanoresonator and have conducted spectroscopy measurements. We have proposed the theoretical interpretation of the observed spectral features of the nonlinear signal generation based on the calculations of eigenmodes of the nanoparticle excited at the fundamental and double frequencies. Our findings suggest the potential of subwavelength semiconductor nanoresonators to become the basis of advanced nanophotonic devices.

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