Ultrafast Optical Modulation of Second- and Third-Harmonic Generation from Cut-Disk-Based Metasurfaces

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ABSTRACT: We design and fabricate a metasurface composed of gold cut-disk resonators that exhibits a strong coherent nonlinear response. We experimentally demonstrate all-optical modulation of both second- and third-harmonic signals on a subpicosecond time scale. Pump–probe experiments and numerical models show that the observed effects are due to the ultrafast response of the electronic excitations in the metal under external illumination. These effects pave the way for the development of novel active nonlinear metasurfaces with controllable and switchable coherent nonlinear response.

KEYWORDS: nanophotonics, plasmonics, nonlinear optics, ultrafast optics, metasurfaces

Metamaterials are artificial materials, assemblies of individual subwavelength units, which can exhibit properties not found in conventional materials. The archetypal example in optics is negative refraction,1 but many other original effects have been proposed and demonstrated, including perfectly focusing superlenses,2 subdiffraction imaging systems,3 cloaking4 and enhanced nonlinearities.5,6 Materials with controllable optical properties are sought after in nanophotonics, as they provide an avenue for designed, ultrafast control of light at the nanoscale,7 leading to signal-processing systems with higher power efficiency, greater bandwidth, and faster response than semiconductor-based electronics allow.8 Applications include all-optical circuits,9 optical memory materials,10 and digital metamaterials, which could lead to programmable all-optical systems.11

Typically weak nonlinear phenomena can be enhanced with plasmonic nanostructures utilizing field enhancement effects.12 Nanoscale plasmonic structures exhibit “hotspots”, which concentrate the energy carried by electromagnetic waves down to nanoscale volumes, providing a way to greatly enhance nonlinear generation efficiencies while also enabling extensive dispersion manipulation.9

Optical metasurfaces,13 two-dimensional metamaterials of subwavelength thickness, are particularly well suited for light control in planar optical elements, with the potential to lead to compact, integrated nanodevices. Their thickness is a small fraction of that of any conventional optical component.14 Although they are most commonly fabricated on thick substrates, flexible supports or membranes have also been used as substrates for metasurfaces.15,16 Moreover, novel fabrication techniques have been developed16–18 that are faster and cheaper than conventional electron-beam lithography, bringing large-scale production closer to practical requirements. Metasurfaces can control beam direction,19–21 shape,22 angular momentum,23 polarization,24,25 and both phase and polarization26 and have been used for lensing27 and holography.28

Optical switching has been demonstrated with metasurfaces in both electro-optical29,30 and all-optical configurations.6,30,31 In the latter case, a metasurface’s optical response is affected by an optical control beam, whose effects can be measured, for example, as a change in the transmission or absorption of a probe beam. In these studies, subpicosecond switching times have been achieved in the extinction of metasurfaces.

Metasurfaces have also been studied in coherent nonlinear optics for the efficient generation of second-33 and third-harmonic signals.34 In particular, since the thickness L of metasurfaces is subwavelength, the interaction length is typically much smaller than the coherence length, Lcoh ≈ 2/Δk,35 and, therefore, phase matching is not a limiting factor for frequency mixing efficiency. Nonlinear planar metasurfaces may be suitable for applications in lasers and optical amplifiers, removing the need for expensive, high-grade macroscopic crystals, which require tedious alignment and millimeter-scale propagation distances. They would enable greater integration and increased portability. Frequency conversion can also be augmented with, for example, steering and switching, to create ultrafast, microscale nonlinear generators with integrated basic signal-processing capabilities, directional control, or both.
In this article, we combine efficient nonlinear generation with dynamic beam modulation on a single metasurface. We experimentally demonstrate ultrafast, all-optical modulation of both second-harmonic generation (SHG) and third-harmonic generation (THG) from nanoscale cut-disk resonator arrays. We explore the mechanisms governing the ultrafast coherent response of the metasurfaces by numerically modeling the time-dependent permittivity of the metal within the random phase approximation (RPA). Our results have important implications for the design of novel metasurfaces with coherent nonlinear response switchable on the picosecond time scale.

■ RESULTS

We studied metasurfaces consisting of arrays of gold nanoscale resonators. Gold (Au), widely used in nanophotonics, is centrosymmetric, prohibiting bulk SHG in the dipolar approximation. The symmetry of the Au crystal lattice is broken at the surface, so that surface effects dominate the second-order nonlinear response, making Au nanostructures possible SH sources.36 A metallic split-ring resonator (SRR) is widely used as the unit cell for metasurfaces in the optical and near-IR spectral ranges, where it exhibits multiresonant behavior.37 Moreover, SRR arrays show strong polarization-dependent second-33 and third-harmonic34 generation. The unit cell of the metasurfaces studied here (Figure 1a) contains a Au disk, 250 nm in diameter and 100 nm in height, with a 140 nm long and 30 nm wide slit cut partway through it. Figure 1b shows a typical SEM micrograph of a 2 × 2 subunit of the array. Au nanodisks are 250 nm in diameter and 100 nm high. They are cut partway with a slit 30 nm wide and 140 nm long and form 60 × 60 µm² arrays. (c, d) Experimental extinction spectra for light polarized (c) along and (d) across the slit. The orientation of the incident wave’s electric field is indicated in the insets.

![Cut-disk metasurface and linear optical properties](image)

Figure 1. Cut-disk metasurface and linear optical properties. (a) Unit cell geometry. (b) Low-voltage SEM micrograph of a 2 × 2 subunit of the array. Au nanodisks are 250 nm in diameter and 100 nm high. They are cut partway with a slit 30 nm wide and 140 nm long and form 60 × 60 µm² arrays. (c, d) Experimental extinction spectra for light polarized (c) along and (d) across the slit. The orientation of the incident wave’s electric field is indicated in the insets.

Au disk, 250 nm in diameter and 100 nm in height, with a 140 nm long and 30 nm wide slit cut partway through it. Figure 1b shows a typical SEM micrograph of a 2 × 2 subunit of the array. In this geometry, the symmetry of a disk-shaped nanoparticle is broken, enabling the linear multiresonant response of the unit cell, as well as its unique coherent nonlinear behavior.

### Linear Optical Properties

To first approximation, a disk supports two nondegenerate dipolar electric resonances, with the dipolar moment either along or perpendicular to the axis of the disk. Given that the distance between neighboring particles is relatively large when compared to the particle’s modal volume, these arrays can be described within the weak coupling regime as the coherent response of isolated resonators. For normally incident light along the z-axis, a uniform disk-shaped particle exhibits a single dipolar resonance, which is degenerate with polarization. This dipolar response is drastically modified when a slit is introduced along the diametric axis of the disks. Two different polarization-sensitive optical responses arise, one with the electric field of the incident wave along the slit ("y" direction in Figure 1c) and the other with the electric field across it ("x" direction in Figure 1d).

In the former case, the extinction spectrum is similar to that of disk-shaped particles, with a single dipolar resonance around 765 nm. In the latter case, the resonance in the same spectral range is blue-shifted by about 90 nm and assumes a double-peak Fano shape. This shape is due to the interference between the dipolar and the quadrupolar modes supported by the disk-shaped particle. The quadrupolar response is dark in a symmetric disk-shaped particle, but the introduction of the slit creates an imbalance in the quadrupolar moment, leading to the appearance of a net dipole moment, making it accessible to incident light through dipolar coupling. The resulting superposition of the spectrally overlapping disk-shaped modes causes an apparent decrease in the extinction of the cut-disk nanoparticle response at frequencies where the two moments interfere destructively. For this geometry the dipole and quadrupole modes of the disk-shaped particle are mixed and form a Fano-type mode. The introduction of the slit also leads to the appearance of a new resonance in the near IR at around 1400 nm. This resonance, also present in square SRRs, is often associated with a magnetic dipole oscillating along the light propagation direction (z-axis) and created by the electrically driven charge oscillations of a solenoid-like current across the particle, resulting in a charge buildup of opposite sign on opposite sides of the slit.58 This resonance provides the enhancement of the electromagnetic field for the generation of coherent nonlinear signals.

### Transient Linear Optical Properties

Upon excitation with ultrashort optical pulses, the linear optical properties of the metasurface and, as a result, its coherent nonlinear response can be modulated via a transient nonequilibrium Au electron energy distribution, resulting in a change in the metal’s permittivity.5 The transient differential extinction (ΔOD) of the metasurfaces (Figure 2a; see Methods for experimental details) reveals a sequence of alternating bands of induced extinction (positive ΔOD) and induced transparency (negative ΔOD) occurring at different wavelengths and time scales. The simulated transient ΔOD spectra (Figure 2b,c; see Methods for the details of the simulations) clearly recover the main features of the experiment. The ground-state simulated extinction (Figure 2d, solid blue line) reproduces the features of the experimental transient extinction (Figure 2f), with the presence of the Fano resonance at visible wavelengths and the magnetic dipole resonance in the near-IR range. Small deviations in the optical response of the simulated and measured structures in both the spectral position and linewidth of the resonances are due to variations in shape and size between particles that affect the position and lead to inhomogeneous broadening of the resonance.40,41

The excited state extinction (Figure 2d, dashed line) and differential extinction (Figure 2e), at a time at which maximum optical modulation is achieved, show a magnetic dipole resonance at around 1400 nm with a clear dispersive behavior,
mainly governed by a combined red-shift and an increased damping of the resonance following optical excitation. The transient response of the Fano resonance is more complex: because of an increased damping for the underlying dipolar mode and radiative loss for the quadrupolar mode, the resonance becomes broader, leading to a loss of coherence between the two resonances and a shallower Fano dip. In the intermediate spectral range (800 to 1200 nm), an induced extinction is observed, mainly due to the broadening of both electric and magnetic resonances. The extinction corresponding to the magnetic resonance reveals the relatively stronger sensitivity of the field associated with the magnetic resonance to transient changes in the particle’s permittivity. This observation is key to understanding the resulting transient coherent response outlined below.

**Coherent Nonlinear Optical Properties.** The cut-disk array behaves as an assembly of in-phase resonators, coherently interacting so as to produce a nonlinear far-field emission diagram with a circular angular spread dependent upon the numerical aperture (NA) of the illumination objective as shown in transmission and reflection measurements (Figure 3a). The SHG spectrum (Figure 3b) shows a double-resonant behavior, indicating the importance of the resonances at both fundamental and SH wavelengths and recovering qualitatively the shape of the Fano resonance. Both SHG and THG signals have maximum emission for an incident polarization across the slit (Figure 3c). In fact, the outgoing intensities of SHG and THG follow a \( \sin^2(\theta_p) \) and \( \sin^4(\theta_p) \) trend, respectively, where \( \theta_p \) is the polarization angle of the fundamental light with respect to the y-axis, with at least a two-order of magnitude difference between minima and maxima. This is due to the \( \sin^2(\theta_p) \) and \( \sin^4(\theta_p) \) dependence of the fundamental light intensity in the respective nonlinear polarizations. We therefore used the fundamental light polarized across the slit (x-polarization) in all the following measurements.

The emitted SHG and THG signals have orthogonal linear polarizations, with THG polarized across the slit, the same as the polarization of the incident fundamental beam, and SHG polarized along the slit, i.e., rotated 90° with respect to the polarization of the fundamental light (Figure 3d). Thus, the dominant tensor component for the effective second- and third-order nonlinear susceptibilities are \( \chi^{(2)}_{xy} \) and \( \chi^{(3)}_{xxx} \), respectively, where the first index refers to the output polarization and the remaining indices describe the polarization of the input beams, i.e., the degenerate fundamental beam in this instance.

**All-Optical Modulation of Nonlinear Processes.** Ultrafast pump–probe experiments were performed to study the transient response of the coherently generated nonlinear signals. The experimental setup (Figure 4a, details in Methods) uses a two-color 200 fs pulsed excitation with a 1400 nm fundamental probe beam (\( \omega_1 \)) and a 1028 nm control light (\( \omega_2 \)). The relative time delay (\( \Delta t \)) between the pulses is scanned over a range of several picoseconds. The spectrum of the nonlinear response of the metasurface obtained for \( \Delta t = 0 \),
when fundamental and control beams are superimposed both spatially and temporally, reveals a series of nonlinear processes (Figure 4a).

When the fundamental and control pulses are not overlapping in time ($\Delta t \rightarrow \pm \infty$), only nonlinear harmonics from the fundamental and control beams ($\omega_1, \omega_2$) are detected. These include the 700 nm SHG1 and 467 nm THG1 from the 1400 nm probe beam as well as the 1400 nm SHG2 from the 1028 nm control beam. As the temporal overlap between the beams is achieved ($\Delta t \rightarrow 0$), two more coherent signals, corresponding to sum-frequency generation (SFG, $\omega_1 + \omega_2$) and four-wave mixing (FWM, $2\omega_2 - \omega_1$), are also observed (Figure 4b). These two signals, therefore, offer a convenient synchronization fingerprint. The dynamics of the single-beam coherent response reveal two differing behaviors. The SHG1 at 700 nm (Figure 4c) experiences a 10% decrease in intensity at $\Delta t \approx 0$, induced by the control beam. A similar behavior is observed for the 467 nm THG1 (Figure 4d), with both signals recovering their steady-state intensity over a time scale of several tens of picoseconds. The temporal response is consistent with an optically induced change in the metal’s permittivity recovery through electron–phonon and phonon–phonon relaxation processes. In contrast, the transient SHG2 excited by the control beam itself at 514 nm (Figure 4e) is enhanced, with a 20% increase in intensity and the relaxation at a time scale of roughly 0.4 ps, much faster than the 700 nm SHG1.

**Analysis and Discussion.** The transient signals of Figure 4c–e are governed by the transient dependencies of both the nonlinear susceptibility of the respective coherent processes and the local amplitude of the fundamental field driving them. This is explicit from the expressions of the nonlinear polarizations $P^{(3)}(2\omega) = \varepsilon_0 \chi^{(3)}(2\omega, \omega_1, \omega_2) E_0^2(\omega)$ for SHG and $P^{(3)}(3\omega) = \varepsilon_0 \chi^{(3)}(3\omega, \omega_1, \omega_2) E_0(\omega)$ for THG, where $E_0(\omega)$ is the local driving field at the fundamental frequency, with $\alpha(\omega)$ being the cut-off linear polarizability. While both $\partial E/\partial E_0$ and $\partial E/\partial E_0$, where $E_0$ is the control field, contribute to the transient response, it is clear from the hierarchy of nonlinear susceptibilities and numerical modeling that the local driving field modification under the influence of the control light is dominating the transient behavior of the nonlinear signals. As one can see from Figure 4d, the polarizability of the cut disk at a wavelength of 1400 nm is significantly decreased under the control illumination due to the damping and red-shift of the magnetic resonance, leading to a decrease in the SHG amplitude around $\Delta t \approx 0$ (Figure 4e). The relatively slow recovery of the signal reflects the resonant behavior of the particle at 1400 nm. A similar argument can be made to explain the transient THG signal of Figure 4d, although, a priori, there is a noticeable difference between the two processes since the SHG at 700 nm is doubly resonant at both the fundamental and SH frequencies, while the THG process is resonant only at the fundamental frequency. The fact that a similar transient response is evident for both signals strengthens the argument that the main mechanism controlling both the decrease and the recovery dynamics of the processes is the ultrafast transient behavior of the local field at the fundamental frequency.

This analysis is further supported by the transient response of the 514 nm SHG2 signal in Figure 4e, showing the SH signal from the control beam at 1028 nm modulated by the simultaneous presence of the probe beam at 1400 nm. This is evidence that, regardless of the intended assignment of control and probe, both beams affect the permittivity of the plasmonic resonators, and their associated nonlinear coherent signals probe those changes. In Figure 4e, both the fundamental and SHG light are nonresonant with the structure, showing a dynamics with an increase in the 514 nm signal followed by a rapid decay to the $\Delta t \rightarrow \pm \infty$ level. The mirror symmetry of the transient response of the 700 nm and 514 nm SHG signals with respect to $t = 0$ is due to the relative arrival time of the corresponding fundamental beams on the sample, with the 1028 nm beam reaching the sample before the 1400 nm beam. The sub-picosecond relaxation time of the 514 nm SHG2 signal is faster than the multi-picosecond relaxation time of the 700 nm SHG1 (Figure 4c,e). This is related to the faster relaxation of the nonlinear response observed in the linear transient spectra (Figure 2b) at the off-resonance wavelength of 1028 nm than at the resonant wavelength of 1400 nm.

**Conclusions**

We studied the ultrafast transient behavior of multiresonant plasmonic nanostructures acting as nonlinear coherent emitters. SHG and THG signals emitted by the cut-disk resonators for both resonant and nonresonant input and output frequencies have been shown to experience an ultrafast modulation of their intensity at the picosecond time scale under optical control. The transient incoherent response of the resonators provided a rationale for these observations in both the sign of the intensity modulation of the coherent nonlinear signals and the respective
relaxation times. In particular, we found that the transient response is governed by the ultrafast response of the electronic density in the metal and its impact on the optical properties of the plasmonic resonators at the fundamental frequency. These results show frequency conversion and its optical nonlinear control combining coherent and Kerr-type nonlinearities of plasmonic nanostructures. As the metasurfaces show directional emission typical of phase-locked resonators, the observed effects can be used for beam steering and directionality control of generated harmonics to allow output harmonic separation with collinear single beam inputs. Further extension of this work may include controlled beam shaping to achieve switchable wavefront generation of nonlinear harmonics.

METHODS

Fabrication. Gold nanostructures were fabricated on a glass substrate via electron beam lithography (EBL) and plasma etching. Exposed negative resist ma-N 2400 was used as the etching mask. After the etching, the residual was cleared with acetone followed by oxygen plasma cleaning. To guarantee the structural stability of the nanostructures, a 5 nm Ta2O5 adhesion layer was deposited by sputtering on the substrate prior to metal lithography. Several arrays were fabricated, on the same sample, with slightly different exposure dose and geometry. For all measurements shown, the array period is 400 nm and the area is $60 \times 60 \, \mu m^2$. AFM measurements show a typical surface roughness of 10%. SEM measurements required low accelerating voltages (2 kV) because the substrate is poorly conductive.

Measurements. For linear transmission measurements, a halogen lamp was used together with polarization optics and a fiber-coupled spectrometer. Extinction is defined as $OD = \frac{-\log_{10}(T/T_0)}{T}$, where $T$ is the transmittance of the array and $T_0$ is a reference transmittance measured through the substrate alone. Raw extinction spectra show uniform 10% noise, which was removed through LOESS interpolation. The setup for nonlinear transmission and pump–probe experiments (Figure 4a) is built around an ultrashort laser source consisting of an amplified ultrafast oscillator (Pharos, Light Conversion) delivering 200 fs pulses at a central wavelength of 1028 nm. This amplified output beam is split in two arms, with roughly half the original 15 W power pumping an OPA (Orpheus, Light Conversion), which generates a frequency-tunable beam. The remaining half of the 1028 nm beam is sent to the sample via a delay line (Thorlabs) that controls the relative arrival time $\Delta t$ of the beams onto the sample. Suitable optical components placed in both beam paths allow for independent control of polarization, power, divergence, and size of the two beams before they are focused onto the sample. For the SH measurements, the fundamental peak irradiance at the sample surface was about 4 GW/cm$^2$, while it was about 2 GW/cm$^2$ for both beams during pump–probe experiments. The laser-induced damage threshold of the samples was observed at peak irradiances of about 10 GW/cm$^2$. The residues of the fundamental incident beams are filtered out after the sample using band-pass filters (Thorlabs). For SHG spectroscopy, the OPA output was tuned to the desired fundamental frequencies. The transient differential extinction measurements were performed using a 100 fs amplified Ti:sapphire laser (Spectra Physics) to generate a 467 nm pump through an OPA (Light Conversion) and a white light continuum (approximately 470–760 nm wavelength range) probe generated using a sapphire crystal. A spatial filter in Fourier space was used to filter out the pump beam and collect the transmitted probe only.

Simulations. The FDTD simulations of Figure 2b–f use Lumerical’s FDTD Solutions package. The simulated structure’s dimensions were taken to be the average sizes obtained through SEM and AFM measurements, resulting in a close match between the extinction peak positions between simulation and linear transmission measurement. However, structure-by-structure variability and fabrication imperfections cause slight shifts and broader peaks in the real case. This does not affect the general behavior induced in the all-optical modulation experiments. The extinction modulation simulations use uniform, temperature-dependent permittivity obtained with a method previously described in detail. In such a model, electron–electron scattering occurring at very short time scales of tens of femtoseconds is neglected, while the experiments do not have the time resolution required to observe these effects.

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All authors have given approval to the final version of the manuscript.

Notes

The authors declare no competing financial interest.

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