Topological insulator metamaterial with giant circular photogalvanic effect

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One of the most notable manifestations of electronic properties of topological insulators is the dependence of the photocurrent direction on the helicity of circularly polarized optical excitation. The helicity-dependent photocurrents, underpinned by spin-momentum locking of surface Dirac electrons, are weak and easily overshadowed by bulk contributions. Here, we show that the chiral response can be enhanced by nanostructuring. The tight confinement of electromagnetic fields in the resonant nanostructure enhances the photoexcitation of spin-polarized surface states of topological insulator Bi1.5Sb0.5Te1.8Se1.2, leading to an 11-fold increase of the circular photogalvanic effect and a previously unobserved photocurrent dichroism ($\Delta_{\text{circ}} = 0.87$) at room temperature. The control of spin transport in topological materials by structural design is a previously unrecognized ability of metamaterials that bridges the gap between nanophotonics and spin electronics, providing opportunities for developing polarization-sensitive photodetectors.

INTRODUCTION

Chirality, or the difference of an object from its mirror image, is a ubiquitous and fascinating phenomenon in nature. It manifests itself at a variety of scales and forms, from galaxies to nanotubes and from organic molecules to inorganic compounds. Detection of chirality at the molecular or atomic level is the key to fundamental sciences (e.g., chemistry, biology, and crystallography) and practical applications (e.g., food and pharmaceutical industry) yet very challenging. Detection of chirality relies on the interaction with electromagnetic fields, which is hindered by the large mismatch between the wavelength of light and the size of most molecules and crystal unit cells, thereby resulting in nearly imperceptible twists of the light field over nanoscale dimensions.

In recent years, the chiral response of topological materials, such as topological insulators (1–4), Dirac/Weyl semimetals (5–8), and two-dimensional (2D) van der Waals heterostructures (9, 10), underpinned by the peculiar quantum features of their electronic structures, have been the subject of intense investigation. For instance, the surfaces of 3D topological insulators support chiral spin currents, a net directional flow of spins in the absence of a net flow of charges. While the spin texture of surface electrons in topological insulators can be unraveled by scanning tunneling spectroscopy (11, 12) or angular resolved photoemission spectroscopy (1, 13, 14), direct electrical or optical addressing of chiral surface states in this class of materials is hampered by the large contribution of the semiconducting bulk to the conductivity, induced by unintentional doping (15–18).

One of the most notable manifestations of the electronic chirality of topological insulator surface states is the dependence of their photocurrent on light helicity. Spin currents can be transformed into polarized net electrical currents when the system is optically driven out of equilibrium (2, 19) by the helicity of light excitation: The absorption of circularly polarized light incident on the surface at oblique angles induces an unbalance between surface carriers of opposite spin at specific points in the $k$-space, thereby generating topological, spin-momentum–locked photocurrents.

Attempts to magnify helicity-dependent photocurrent (HDPC) in topological materials have relied on strategies to increase the inherent surface-to-bulk contribution, such as using Bi-chalcogenides with low intrinsic doping (3, 20–22), reducing the crystal thickness to only a few quintuple layers (23), tuning the Fermi energy by electrical gating (24), or selectively exciting surface carriers below the bulk bandgap (25). More recently, research has broadened to Weyl semimetals, where the “chiral anomaly” (6, 7, 26), induced by unpaired Dirac cones, is expected to yield highly chiral photocurrents.

Designer metamaterials, with structural features comparable to the wavelength of light, provide an independent approach to devise optical properties on demand and enhance light-matter interaction (27). The creation and enhancement of optical chirality by metamaterials is particularly interesting and elaborate (28–31). For instance, patterning an achiral medium (e.g., a metal) with chiral, subwavelength shapes makes the resulting metamaterial intrinsically chiral. Similarly, the mutual orientation of a metamaterial, patterned with nonchiral structures, and the incident electromagnetic wave can lead to an extrinsically chiral response (32, 33). Both forms of chirality can emerge in 2D and 3D metamaterial systems and lead to strong optical activity, circular dichroism, and asymmetric transmission.

Here, we provide a first demonstration of the use of artificial nanostructuring to enhance the chiral photogalvanic response of topological insulators. Thanks to the tight confinement of electromagnetic fields, resonant nonchiral metamaterials effectively enhance the photoexcitation of spin-polarized states, thereby increasing the unbalance between surface-state carriers with opposite spin helicity and overall resulting in a giant enhancement of the extrinsic chiral photocurrent response of a 3D topological insulator (TI), Bi1.5Sb0.5Te1.8Se1.2 (BSTS).
RESULTS AND DISCUSSION

Helicity-dependent photocurrent

Surface carriers can be selectively excited in topological insulators by circularly polarized light directed at oblique incidence on the surface of the crystal, and the resulting current flow is determined by the spin-momentum locking of the carriers (Fig. 1). As the topological insulator crystal is intrinsically achiral (the surface states have equal number of carriers with opposite spin orientations), and the relevant spins lay in the surface plane of the crystal, photocurrent at normal incidence does not generate any HDPC. However, spin-selective photoexcitation of surface-state carriers by obliquely incident light with a given helicity induces chirality (unbalance in the populations of surface state carriers with opposite spin orientations), akin to the extrinsic chirality described for metamaterials. This effect, known as the circular photogalvanic effect (CPGE) (34–36), is illustrated in Fig. 1A, where a light beam of defined handedness selectively promotes carriers of matching spin from surface to bulk conduction bands, thus leaving the surface bands asymmetrically depopulated in k-space: The excess surface carriers of opposite spin generate the net spin-polarized electrical current, \( j_{\text{s}} \). The presence of a nanostructure on the surface of topological insulators, designed to have a resonant absorption at the wavelength of excitation, effectively increases the number of surface conduction carriers that are promoted to the bulk conduction bands (Fig. 1B), thus significantly enhancing the CPGE contribution to the photocurrent. Figure 1C shows the experimental arrangement to measure HDPC: Light is incident at a variable angle, \( \theta \), on the surface of the topological insulator crystal (xy plane), and the light polarization is changed continuously from linear to circular by rotating a quarter-wave plate, whose fast axis forms an angle, \( \phi \), with the polarization axis. Without any applied bias, an HDPC flows across the two gold contacts on the crystal surface.

Optical absorption and metamaterial design

To assess the enhancement of chirality exclusive to topological surface spin currents, we chose a metamaterial design that does not introduce optical chirality, neither intrinsic nor extrinsic (37). The metamaterial unit cells consist of square rings of \(~200\text{-nm}\) lateral size and \(~100\text{-nm}\) ring width, arranged on a square lattice of \(~335\text{-nm}\) period. Such pattern was carved by focused ion beam (FIB) milling between two gold electrodes deposited on a ~250-nm-thin BSTS flake, as shown in the scanning electron microscopy (SEM) images in Fig. 1D.

The metamaterial was designed to resonantly increase the optical absorption of the unstructured BSTS flake at the excitation wavelength, \( \lambda = 532 \text{ nm} \), at both normal (\( \theta = 0^\circ \)) and oblique (\( \theta = 45^\circ \)) incidence (Fig. 2). The experimental and numerically simulated spectra of the BSTS metamaterial (Fig. 2A, right) show a resonant increase in optical absorption with respect to the case of unstructured BSTS (Fig. 2A, left). Following from the design symmetry, the metamaterial geometry does not induce any helicity dependence to the bulk response of the BSTS flake, as confirmed by the maps of electric field intensity for circularly polarized light of opposite handedness (Fig. 2, B and C).

Control of CPGE in topological insulator by the metamaterials

We measured HDPC under nearly uniform illumination (laser beam diameter of ~200 \( \mu \text{m} \), much larger than the BSTS device size of ~10 \( \mu \text{m} \)), with no applied bias (Fig. 3). The residual un-uniformity of illumination results in thermal gradient that induces polarization-independent photothermoelectric currents, which contribute to the photocurrent background. This component of the current is sensitive to the position of the excitation beam on the sample (3).

To measure a clearly distinguishable photocurrent signal, we adjusted the laser beam position on the BSTS flake and BSTS metamaterials to obtain the maximum total photocurrent, i.e., we maximized the thermoelectric photocurrent offset (see section S1 and figs. S1A and S2). Contributions of surface carriers to the photocurrent, seen in HDPC, are allowed only at oblique illumination (2), as illustrated in the top section of Fig. 3A. We measured the photocurrent, \( j_{\text{a}} \), shining the laser at \( \theta = -45^\circ \) and going through a full polarization cycle of its polarization angle, \( \phi \), from \( 0^\circ \) to \( 360^\circ \). This induces a continuous change of the incident polarization, from linear (\( \phi = 0^\circ \)), \( 90^\circ \) to circular of right (\( \phi = 45^\circ \)) and left (\( \phi = 135^\circ \)) handedness, with \( 180^\circ \) period.

The bottom panel of Fig. 3A shows the photocurrent, \( j_{\text{a}} \), measured in an unstructured BSTS flake (black-filled squares). The current has the characteristic polarization-dependent behavior observed in other Bi-chalcogenide topological insulators (3, 20, 24) and other 2D material systems [i.e., quantum wells (34, 35), transition metal dichalcogenides (8, 38, 39), and Weyl semimetals (6, 7, 26)]. It

![Fig. 1. HDPC in topological insulators and topological insulator metamaterials.](Image)
composes of four components, expressed by the phenomenological equation (3, 35)

\[ j_a = C \sin 2\phi + L_1 \sin 4\phi + L_2 \cos 4\phi + D \]  

(1)

The coefficients \( D \) and \( L_2 \) are associated to photocurrent contributions from the semiconducting bulk. Specifically, \( D \) is related to the polarization-independent photothermal electric background current, which sets the overall directional current flow, while \( L_2 \) to the photon drag effect, which results from linear momentum transfer of the incident photons to the excited carriers. Conversely, the coefficients \( L_1 \) and \( C \) quantify photocurrent contributions from surface carriers. Such currents, driven by linear and circular polarization of the incident light, originate from the linear and circular photogalvanic effects, respectively. The dependence of the photocurrent on helicity is seen in the small asymmetry between the peaks corresponding to photoexcitation by light of opposite handedness (\( \sigma^+ \) and \( \sigma^- \)) in the bottom panel of Fig. 3A. The relative contribution of each surface

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Fig. 2. Optical absorption enhancement in BSTS achiral metamaterial. (A) Measured (dashed lines) and simulated (continuous lines) optical absorption of an unstructured BSTS flake and a nanostructured BSTS metamaterial (experimental data were collected with unpolarized light at normal incidence and numerical aperture = 0.7, while simulations correspond to circularly polarized light incident at \( \theta = 0^\circ \) and \( \theta = 45^\circ \)); at \( \lambda = 532 \) nm, the absorption of the BSTS metamaterial is \( \sim 0.7 \), twice that of the unstructured BSTS flake \( \sim 0.35 \). (B and C) Maps of the electric field intensity, \(|E|^2\), at 10 nm below the top surface of the metamaterial unit cell at normal incidence, for left (LCP) and right (RCP) circular polarization, respectively.

Fig. 3. Multifold increase of CPGE in BSTS topological insulator by metamaterials. (A) (top) Schematic of HDPC in an unstructured BSTS flake; (bottom) experimental photocurrent measured on an unstructured BSTS flake, at room temperature, and fitting with Eq. 1, showing the expected 4\( \phi \) dependence and a small 2\( \phi \) asymmetry between right (\( \sigma^+ \)) and left (\( \sigma^- \)) circularly polarized illumination; (inset) fitting coefficient \( C, L_1, L_2, \) and \( D \), indicating a predominance of bulk photon-drag contribution, \( L_2 \), in the photocurrent modulation. (B) (top) Schematic of light HDPC in a nanostructured BSTS metamaterial; (bottom) experimental photocurrent measured on a nanostructured BSTS metamaterial, at room temperature, and fitting with Eq. 1, showing how the metamaterial induces a 2\( \phi \) dependence that indicates that the sample responds almost exclusively to right (\( \sigma^+ \)) and left (\( \sigma^- \)) circularly polarized illumination; (inset) fitting coefficient \( C, L_1, L_2, \) and \( D \), indicating a predominance of CPGE, \( C \), in the photocurrent modulation.
and bulk component to the total photocurrent is shown in the bar plot of the coefficients of Eq. 1 (inset of Fig. 3A). While surface contributions to the photocurrent are discernible even at room temperature in the unstructured BSTS samples [thanks to the large surface-to-bulk conductivity known for this particular stoichiometry (22, 40–42)], their dependence on light helicity is overshadowed by bulk components (\(C/D = 0.03\)) and too small for any practical device or application.

The very same BSTS flake, patterned with square ring metamaterial array, behaves markedly differently. The resonant metamaterial structure induces much larger asymmetry in the population of surface conducting bands, increasing the net spin current (top schematic in Fig. 3B). The effect is so strong that the measured photocurrent (black-filled circles in bottom panel of Fig. 3B) has a distinct 2\(\varphi\) evolution as function of polarization, irrespective of the contribution of linearly polarized components (43). The effect is even more apparent when comparing the coefficients of Eq. 1 (inset of Fig. 3B), where, now, \(C\) is far higher than the linear coefficients, \(L_1\) and \(L_2\), and its value accounts for a significant fraction of the photocurrent (\(C/D = 0.33\)). Table 1 summarizes the fitting coefficients of photocurrents, \(j_\sigma\), for both unstructured and nanostructured BSTS, at and off normal incidence. The photocurrent circular dichroism induced by spin-polarized surface states can be defined as (35)

\[
\rho_{\text{circ}} = \frac{|I_\sigma^+ - I_\sigma^-|}{|I_\sigma^+ + I_\sigma^-|}
\]

where \(I_\sigma^+\) and \(-I_\sigma^-\) are, respectively, the values of photocurrents under left and right circularly polarized optical excitation, excluding the spin-insensitive component \(D\) (44). A threefold increase of the degree of spin polarization of unstructured BSTS (\(\rho_{\text{circ}} = 0.26\)) is seen in the BSTS metamaterial (\(\rho_{\text{circ}} = 0.87\)), an unprecedented degree of spin polarization approaching unity even at room temperature in nonmagnetic materials (44–47). Additional validation of the functional dependence of the HDPC on incidence angle \(\theta\) is shown in the Supplementary Materials (section S2 and figs. S3 and S4). Notably, the surface nature of HDPC in BSTS was also confirmed, repeating the experiments on a trivial chalcogenide insulator crystal (Fig. 4). Here, we describe the unstructured BSTS crystal by its isotropic relative permittivity, \(\epsilon_r\) (experimentally determined by ellipsometric measurements (49)), modified by ad hoc off-diagonal terms of the permittivity tensor to mimic the effective optical chirality induced by the in-plane spin of Dirac surface electrons. We performed full-wave electromagnetic simulations for both unstructured and nanostructured BSTS, replicating the sample illumination conditions used in the experiments (\(\lambda = 532\) nm; \(\theta = 0, \pm 45^\circ; \varphi = 0^\circ\) to 360°) and integrated the electric field intensity within the top 3 nm (50) to evaluate the optical absorption at the surface of the topological insulator.

The electromagnetic modeling (Fig. 4B) is in excellent agreement with the experimental HDPC result (Fig. 4A), corroborating the link between anisotropic optical absorption at the BSTS surface and selective spin-photocurrent generation. The normalized polar plots in Fig. 4 (A and B) provide an immediate visualization of the prominence of linear components (four lobes) versus circular components (two lobes) in unstructured and nanostructured BSTS. While at normal incidence, the BSTS metamaterial does not produce any notable effect; at oblique incidence, it filters out linear components, distilling the response to circularly polarized light illumination and leading to a giant enhancement of the chiral photocurrent. The degree of chirality predicted by the simulations, according to the same Eq. 2, is \(\rho_{\text{circ}}\) = 0.18 for the unstructured BSTS and \(\rho_{\text{circ}}\) = 0.89 for the BSTS metamaterial, in excellent agreement with the values obtained experimentally (refer also to section S4 and fig. S7). Furthermore, in both experiments and simulations, illumination from mirror directions of incidence (with respect to the normal) yields opposite phases, just as expected for extrinsic chirality.

To conclude, the hitherto unrecognized ability of metamaterials to control surface transport in topological insulator by structural design provides a powerful toolbox to bridge the gap between nanophotonics and spin electronics. We have shown that resonant nanostructures can be used to induce giant enhancement of the extrinsic chiral photocurrent response of a topological insulator. We argue that this approach could be used to exert control over spin-transport properties of other classes of quantum and topological materials.

### Electromagnetic modeling

In the following, we describe the photocurrent behavior of spin-transport metamaterials by electromagnetic modeling. The generated photocurrent is directly proportional to the optical absorption, carrier density, mobility, and lifetime of the topological insulator. Thus, under the assumption that the optical absorption of the BSTS metamaterial increases upon nanostructuring, while the remaining transport parameters remain unaltered, carrier anisotropy can be mapped to an anisotropic optical model of the BSTS topological insulator crystal (Fig. 4). Here, we describe the unstructured BSTS crystal by its isotropic relative permittivity, \(\epsilon_r\), theoretically determined by ellipsometric measurements (49), modified by ad hoc off-diagonal terms of the permittivity tensor to mimic the effective optical chirality induced by the in-plane spin of Dirac surface electrons. We performed full-wave electromagnetic simulations for both unstructured and nanostructured BSTS, replicating the sample illumination conditions used in the experiments (\(\lambda = 532\) nm; \(\theta = 0, \pm 45^\circ; \varphi = 0^\circ\) to 360°) and integrated the electric field intensity within the top 3 nm (50) to evaluate the optical absorption at the surface of the topological insulator.

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### Table 1. Fitting coefficients of the HDPC in unstructured BSTS flake and nanostructured BSTS metamaterials.

| \(\theta (^\circ)\) | BSTS flake | \(C\) (pA) | \(L_1\) (pA) | \(L_2\) (pA) | \(D\) (pA) |
|-----------------|------------|-------------|-------------|-------------|-------------|
| -45             | Unstructured | -1.1 ± 0.1  | 0.6 ± 0.1   | -3.1 ± 0.1  | 35.4 ± 0.1  |
|                 | Nanostructured | -12.7 ± 0.3 | 4 ± 0.3     | -1.8 ± 0.3  | 38.3 ± 0.3  |
| 0               | Unstructured | 0.05 ± 0.05 | -0.002 ± 0.05 | 0.09 ± 0.05 | 8.5 ± 0.05  |
|                 | Nanostructured | -0.2 ± 0.04 | -0.2 ± 0.04 | 0.04 ± 0.04 | 11.4 ± 0.04 |
| 45              | Unstructured | 1.0 ± 0.1   | 0.32 ± 0.1  | -3.2 ± 0.1  | 35.1 ± 0.1  |
|                 | Nanostructured | 11.4 ± 0.4  | -0.9 ± 0.4  | -3 ± 0.4    | 41.8 ± 0.4  |
(e.g., Weyl semimetals and van der Waals heterostructures) and to find application in integrated spin-polarized photodetectors that are in great demand for ultrasensitive chiral molecular sensing and quantum optospintronic devices, where polarization and entanglement could be transferred from photons to electron spins.

**MATERIALS AND METHODS**

**Device fabrication**

Bi$_{1.5}$Sb$_{0.5}$Te$_{1.8}$Se$_{1.2}$ single crystals were grown using a modified Bridgeman method. This particular BSTS stoichiometry yields large surface-to-volume conductivity, so that transport in nanometric thin flakes is surface dominated (22). BSTS flakes were mechanically exfoliated and transferred from the bulk crystals onto a SiO$_2$ (285 nm)/P-Si substrates. Electrical contacts (Cr/Au 5/50 nm) for photocurrent measurements were created on the devices by electron beam lithography and thermal evaporation. Square ring metamaterials were carved on the BSTS flake between the contacts by FIB milling. All fabrication steps were performed minimizing the exposure of BSTS to the electron and ion beams. SEM images were acquired after photocurrent measurements.

**HDPC measurement**

Photocurrent measurements were performed at room temperature, illuminating the devices with continuous wave laser ($\lambda = 532$ nm) and with no applied bias. The linearly polarized laser beam was modulated at a frequency of 137 Hz by an optical chopper before passing through a $\lambda/4$ retarder and focused to a spot size of ~200-$\mu$m diameter between the electrodes. The experimental setup allowed to continuously vary the polarization of incident light from linear (s-polarized) to circular (right circular polarization [RCP] and left circular polarization [LCP]) by rotating the $\lambda/4$ wave plate. The incident light polarization was calibrated by a polarimeter. The photocurrent was measured with a lock-in amplifier referenced to the light modulation frequency. In all samples, the photocurrent was found to be linearly dependent on excitation intensity (refer to fig. S1B). All the measurements were performed at a constant illumination intensity of 10 W cm$^{-2}$.

**Electromagnetic simulations**

3D electromagnetic simulations were performed using COMSOL Multiphysics. The square ring metamaterial dimensions were derived from the SEM images. The relative permittivity of BSTS, $\varepsilon_{r,\text{ell}}$ (isotropic), was experimentally obtained by ellipsometry (see section S5 and fig. S8). The simulations were built to match the experimental arrangement shown in Fig. 1C: The square rings were laying on the $xy$ plane and extruded through the $z$ direction; the direction of incidence of the electromagnetic wave was lying in the $xz$ plane, with an elevation angle, $\theta$ ($0^\circ$, $\pm45^\circ$), with respect to the $z$ axis; the

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**Fig. 4. Distillation of CPGE in BSTS topological insulator flakes by nanostructured metamaterials.** (A) Normalized polar plots of HDPC for an unstructured BSTS flake (left column) and a nanostructured BSTS metamaterial (right column) at three different angles of incidence, $\theta = 45^\circ$ (top row), $\theta = 0^\circ$ (center row), and $\theta = -45^\circ$ (bottom row); at $\theta = 0^\circ$, mostly $L_2$ contributes to the modulation of the photocurrent; at $\theta = 45^\circ$ and $\theta = -45^\circ$, the HDPC patterns of the unstructured BSTS flake and the BSTS metamaterial are distinctly different: The metamaterial distills the $C$ term contribution to the photocurrent modulation, with respect to the unstructured BSTS flake, where $L_1$, $L_2$, and $C$ have comparable amplitudes. (B) Simulated $|E|^2$ in both unstructured 250-nm BSTS film (left column) and nanostructured BSTS metamaterial (right column) at three different angles of incidence, $\theta = 45^\circ$ (top row), $\theta = 0^\circ$ (center row), and $\theta = -45^\circ$ (bottom row), with artificially increased chirality of the optical permittivity, $\varepsilon_{r,\text{ell}}$, of the BSTS; the distinct behavior observed in the unstructured BSTS and the BSTS metamaterial matches remarkably well with the measured photocurrent, indicating how the chirality of the surface carriers and the increased absorption given by the metamaterial result in a giant increase of the CPGE; the $|E|^2$ is integrated in a 3-nm-thin slab at the surface of the film. To better visualize the polarization ($\phi$) dependence of both experimental data and numerical model, we subtract the polarization independent background from each curve and normalize them.
incident wave polarization was obtained by introducing a $\varphi$ dependence to the two orthogonal components ($h, v$) of the electric field with a $\pi/2$ phase retardation: $h = (1 - i)^*|\sin(\varphi)|\cos(\varphi)|$, $v = |\sin(\varphi)|^2 + |\cos(\varphi)|^2$. The simulations of the optical absorption (Fig. 2A) of the square ring BSTS metamaterials were performed using the isotropic, ellipsometrically measured $\varepsilon_{\text{r, ell}}$. The simulations of $|E|^2$ (Fig. 4B and fig. S7) were obtained by assigning nonzero values to the off-diagonal terms of the permittivity tensor. The diagonal terms of the permittivity tensor were defined as $\varepsilon_{xx,yy,zz} = \varepsilon_{r, ell}$, while the off-diagonal terms were defined as $\varepsilon_{yz,zy} = \pm i\chi$, with $\chi = 0.36\varepsilon_{r, ell}$. The electric field intensity $|E|^2$ plotted in the graphs of Fig. 4B and fig. S7 was obtained by performing a volume integration of $|E|^2$ within a 3-nm-thin slab from the surface. The anisotropic response along the $x$ direction was obtained by assigning nonzero values to the off-diagonal terms of the permittivity tensor, $\varepsilon_{r, ell}(x,y,z) = \pm \chi_\ell \varepsilon_{r, ell}(x,y,z)$. The incident wave polarization rotation, reproducing the experimental arrangement of polarizer and quarter-wave plate, was obtained by introducing a $\varphi$ dependence to both $s$ and $p$ components of the electric field with a $\pi/2$ phase retardation between the two. The electric field intensity plotted in the graphs was obtained by integrating $|E|^2$ within a 3-nm-thin slab from the surface.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/14/eabe5748/DC1

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