Surface Polariton-Like s-Polarized Waveguide Modes in Switchable Dielectric Thin Films on Polar Crystals

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Surface phonon polaritons (SPhPs) and surface plasmon polaritons (SPPs), evanescent modes supported by media with negative permittivity, are a fundamental building block of nanophotonics. These modes are unmatched in terms of field enhancement and spatial confinement, and dynamical all-optical control can be achieved, e.g., by employing phase-change materials. However, the excitation of surface polaritons in planar structures is intrinsically limited to p-polarization. On the contrary, waveguide modes in high-permittivity films can couple to both p- and s-polarized light, and in thin films, their confinement can become comparable to surface polaritons. Here, it is demonstrated that the s-polarized waveguide mode in a thin Ge$_3$Sb$_2$Te$_6$ (GST) film features a similar dispersion, confinement, and electric field enhancement as the SPhP mode of the silicon carbide (SiC) substrate, while even expanding the allowed frequency range. Moreover, it is experimentally shown that switching the GST film grants nonvolatile control over the SPhP and the waveguide mode dispersions. An analytical model is provided for the description of the GST/SiC waveguide mode and it is shown that the concept is applicable to the broad variety of polar crystals throughout the infrared spectral range. As such, complementarily to the polarization-limited surface polaritons, the s-polarized waveguide mode constitutes a promising additional building block for nanophotonic applications.

1. Introduction

The field of nanophotonics pursues control over near-field optical phenomena that can be tailored in custom-designed nanoscale material systems. Key to achieve this control are surface-bound modes such as surface plasmon polaritons (SPPs) in metals or surface phonon polaritons (SPhPs) in polar crystals, which stand out by their unprecedented ability to confine and enhance the optical field in subwavelength structures. In the past decades, the potential of these modes has been demonstrated by the large number of applications found in the fields of spectroscopy and sensing, photonic circuitry, lasers, and nonlinear optical phenomena.

SPPs and SPhPs are electromagnetic evanescent modes bound to the interface of two materials with permittivities of opposite sign. SPPs are supported in metals doped semiconductors from infrared (IR) up to visible frequencies, where the negative Re(ε) arises from the Drude response of free electrons, whereas SPhPs arise in the mid to far IR range in the reststrahlen band of polar crystals between the transverse optical (TO) and longitudinal optical (LO) phonon frequencies. In comparison, SPhPs are valued for their longer lifetimes than SPPs due to the low losses of the driving phonon resonances.

Fundamentally, however, both modes consist of collective charge oscillations coupled to light, leading to shared properties such as subwavelength confinement, strong field enhancement, an asymptotic dispersion, and waveguiding characteristics, which are the reasons for the versatile usability of polaritonic modes.

However, SPPs and SPhPs in planar heterostructures intrinsically carry out-of-plane electric fields, and thus can only be excited by p-polarized light, while no s-polarized light can couple to them. This polarization restriction limits the versatility of polariton-based nanophotonics and potentially hinders the development of future technologies, such as emission control or exploiting solar energy. Very recently, omnipolarization waveguide modes in high-permittivity planar media were proposed to circumvent the polarization bottleneck of polaritons, showing that in thin films, waveguide modes can reach comparable degrees of confinement as polaritons. However, while gaining a waveguide mode capable of coupling to s-polarized light, a high-permittivity thin film does not support a SPP or SPhP.

Another drawback of surface polaritons is their lack of active tunability when considered on a plain metal (polar crystal),...
because there the SPP (SPhP) dispersion is bounded by the fixed plasma frequency (optical phonon frequencies). For SPPs, this disadvantage can be overcome in doped semiconductors or graphene, where the photo or voltage-induced free charge carrier concentration enables tuning of the plasma frequency. A similar approach has been proposed for SPhPs but here, the tunability of the SPhPs gained by an additional photoinduced plasma contribution is limited.

A significantly larger and nonversatile switching contrast can be achieved by utilizing phase-change materials (PCMs) such as the Ge_{3}Sb_{2}Te_{5} compound (GST), which can be reversibly switched between its amorphous (a-) and crystalline (c-) phases by ultrafast electrical or optical pulses. The high permittivity contrast between the two phases (\(\varepsilon_{a-GST} \approx 13.8 + 0i\) and \(\varepsilon_{c-GST} \approx 31.4 + 9i\); values are obtained from a global fitting procedure, see the Experimental Section for details) in the 1R is sensed by SPhPs that evanescently penetrate the GST, leading to a significant shift of the SPhP mode frequency upon switching of the GST phase. However, while systems utilizing PCMs have enabled active control over surface polaritons, the large potential of s-polarized waveguide modes in these systems remains, so far, unexploited.

2. Concept

In this work, we suggest a generic, actively tunable material system for the infrared spectral range that simultaneously supports p-polarized SPhPs and s-polarized waveguide modes. Intriguingly, the waveguide modes replicate the properties of the SPhP that define its suitability for nanophotonic applications, such as the enhancement of local electric fields and sub-wavelength confinement. We experimentally demonstrate active tuning of both excitations with exceptionally high tuning figures of merit by means of a phase-change material, which, at the same time, constitutes the waveguide core. As a model system, we analyze the tuning of SPhP and waveguide modes in a GST/SiC structure, where the GST film can be switched between its amorphous and crystalline state. Offering broad functionality, such as polarization dependent polariton optics, the proposed material system features a unique combination of active tunability of its guided modes and the lifting of the polarization bottleneck of conventional polaritons.

An SPhP propagating along the surface of a polar crystal such as SiC exhibits an in-plane \(E_x\) field that peaks at the interface (Figure 1a). We define the \(x-z\) plane as the plane of incidence, and all waves propagate in \(x\)-direction. In Figure 1e (and zoomed into the reststrahlen region of SiC in Figure 1f), the dispersion of this SPhP is shown in black. By placing a 400 nm thick a-GST onto SiC, the SPhP dispersion is red-shifted (blue line in Figure 1e,f), while maintaining the shape of its in-plane electric field \(E_x\) (Figure 1b). Interestingly, here, the maximum field is encountered at the air/a-GST interface, and decays exponentially through the GST film into the SiC substrate (please note that for illustration purposes, the GST film thickness in Figure 1 a–d is not to scale with respect to the shown \(z\)-range of the adjacent air and the substrate).

A freestanding 400 nm thin a-GST film, on the other hand, supports an s-polarized waveguide mode owing to the high index contrast between a-GST and air. The in-plane electric field \(E_y\) of this mode is shown in Figure 1d, and the corresponding dispersion is plotted in green in Figure 1e,f. Notably, this mode features a much larger penetration depth \(\delta\) in positive and negative \(z\)-direction, with \(E(z) = E(0)e^{-z/\delta}\), than the SPhP, hence it is much less confined within the a-GST film. The dispersion of this waveguide mode, however, is steeper and covers a much larger frequency range than the SPhP, which is restricted to the SiC reststrahlen band. Now, the combined system of a-GST/SiC supports not only the shifted p-polarized SPhP mode but simultaneously, at the

![Figure 1. Guided modes in an a-GST film on an SiC substrate.](image-url)

- a–d) Schematic in-plane electric field distributions of SPhPs and waveguide modes, excitable with p- and s-polarized light, respectively, supported a) on a bare SiC substrate, b,c) in a 400 nm thin a-GST film on SiC, and d) in a freestanding a-GST film. e,f) Dispersions of the four modes shown in (a)–(d), where (f) zooms into the SiC reststrahlen region between the TO and LO frequencies (\(\omega_{TO}\) and \(\omega_{LO}\)), marked by the dotted rectangle. The SiC substrate leads to a flattening of the waveguide mode (red line) compared to the air-bound mode (green line), leading to a comparable dispersion to the SPhP (black and blue lines) in the dispersion window shown in (f).
same frequency for smaller in-plane momenta, an s-polarized waveguide mode localized in the a-GST film. In Figure 1c, the in-plane electric field $E_y$ of this air/GST/SiC-bound mode is shown, featuring a higher confinement than the air-bound waveguide mode due to the SiC substrate. Strikingly, the dispersion of the air/GST/SiC-bound waveguide mode (red line in Figure 1e,f), while approaching asymptotically the dispersion of the air-bound waveguide mode at large in-plane momenta $k$, experiences an upward bending for small $k$, becoming quite similar to the dispersion of the SPhP. Thus, the combined structure of a thin a-GST film on a SiC substrate not only supports both p-polarized SPhPs and s-polarized waveguide modes, but thanks to the reststrahlen characteristics\cite{38} of the SiC substrate, the waveguide mode mimics the SPhP for small in-plane momenta $k/k_0$ in both field distribution and dispersion.

An a-GST film of $\approx 400$ nm thickness, as discussed in Figure 1, yields optimal polariton-like characteristics of the waveguide mode, such as its field enhancement (see Figure 5 for details). However, the second aspect of our concept is the active tunability of the GST/SiC system via GST phase switching and the resulting tuning potential for both the SPhP and the waveguide mode. We find that the maximum frequency shift of the SPhP upon switching of the GST phase is obtained for an a-GST film thickness of $\approx 150$ nm (see Figure S1 in the Supporting Information). Importantly, for this film thickness, the SPhP-like dispersion and a significant field enhancement of the waveguide mode are still conserved. Thus, experimentally, we choose to investigate a GST/SiC system with $\approx 150$ nm thickness of the a-GST film.

3. Experiment

A 144 nm thin a-GST film was sputter-deposited onto a 4H-SiC substrate and placed into a home-build Otto-type prism coupling setup, sketched in Figure 2a (see the Experimental Section). In short, the incoming light impinges on the prism surface at an angle $\theta$ above the critical angle of total internal reflection ($\approx 24.6^\circ$ for KRS5). The generated evanescent wave in the air gap between prism and sample couples to the evanescent, bound modes in the sample, resulting in attenuated total reflection dips at the resonances in a reflectance spectrum. By taking several spectra at varying incident angle, dispersions can be mapped out.\cite{39} As an excitation source for the reflectance spectroscopy measurements, we employed a mid-IR free electron laser (FEL) with tunable wavelength in the range of 3-60 $\mu$m and a spectral line width of $\approx 0.3\%$\cite{40} (see the Experimental Section). In a first step, all experimental data were obtained for a-GST, and after crystallizing the a-GST on a heating plate (300°C, 5 min), the measurements were repeated for the c-GST phase. Note that crystallization of a-GST is accompanied by a density shift of the SPhP upon switching of the GST phase is obtained and the waveguide mode. We find that the maximum frequency shift of the SPhP upon switching of the GST phase is obtained for a-GST film thickness of $\approx 150$ nm (see Figure S1 in the Supporting Information). Importantly, for this film thickness, the SPhP-like dispersion and a significant field enhancement of the waveguide mode are still conserved. Thus, experimentally, we choose to investigate a GST/SiC system with $\approx 150$ nm thickness of the a-GST film.

Furthermore, a Lorentzian line shape was fitted to the experimental reflectance dips in order to extract frequency positions $\mu$ and full width at half maxima (FWHM $= 2\sigma$) of the resonances. Interestingly, the measured quality factor ($Q = \mu$/FWHM) in a-GST of the waveguide mode ($Q = 109$) is even higher than that of the SPhP ($Q = 82$), whereas in c-GST, the waveguide mode has a broader resonance ($Q = 14$) than the SPhP ($Q = 27$). For the case of a-GST, these $Q$-factors easily lie in the range of the high-$Q$ resonances arising from localized SPhPs in polar crystal nanostructures (45-360).\cite{43,48} The generally lower $Q$-factors in c-GST, on the other hand, can be attributed to nonzero

**Figure 2.** Tuning of the SPhP and the waveguide mode by switching the GST phase. a) Setup of the prism coupling experiments in the Otto geometry with variable air gap between the KRS5 ($n_{KRS5} = 2.4$) prism and the sample, allowing for momentum-matched, critical coupling to evanescent modes supported by the sample such as SPhPs and waveguide modes. b) Reflectance spectra at $\theta = 30^\circ$ for the crystalline and amorphous GST phases for s- and p-polarized incident light of a 144 nm (132 nm) thin a-GST (c-GST) film on SiC, demonstrating the tuning of both SPhP and waveguide mode upon switching the GST phase. A global fitting algorithm was used to fit simulated reflectances (dashed-dotted lines) to the experimental spectra (solid lines). (blue and green curves) or an s-polarized waveguide mode (red and orange curves). All data were fitted with a global fitting procedure (see the Experimental Section for details), and the resulting theoretical spectra calculated by means of a $4 \times 4$ transfer matrix formalism\cite{43} are also shown in Figure 2b, nicely reproducing the experimental data. (The low amplitude of the waveguide mode in c-GST arises due to a prism coupling gap below the gap of critical coupling,\cite{39} see the Experimental Section for details).
absorption ($\text{Im}(\varepsilon_{\text{c-GST}}) = 9$) compared to a-GST. However, it is noteworthy that the waveguide mode features similar $Q$-factors as the SPhP in both GST phases, providing first evidence that the waveguide mode can indeed be seen as a polariton-like excitation with s-polarization.

Reflectance spectra were taken for incident angles of $26^\circ$–$34^\circ$ in steps of $1^\circ$ resulting in reflectance maps that reproduce the dispersion of the corresponding mode. Maps for bare SiC, a-GST/SiC, and c-GST/SiC for p-polarized incident light are shown in Figure 3a–c, respectively, and maps for a-GST/SiC and c-GST/SiC for s-polarized incident light in Figure 3e,f. In Figure 3d, an overview over all modes is given, and the respective dispersion curves are additionally plotted on top of the individual reflectance maps. The theoretical dispersions were extracted from the imaginary part of the correspondingly polarized reflection coefficient obtained by transfer matrix calculations using the parameters from the global fitting procedure. All experimental dispersions are well reproduced by the simulations.

For 144 nm thin a-GST and at $\theta = 30^\circ$, the SPhP shifts by $\delta \omega = 27 \text{ cm}^{-1}$ upon switching the GST phase, and the waveguide mode by $69 \text{ cm}^{-1}$. This corresponds to a tuning figure of merit (TFOM) of $\text{TFOM} = \delta \omega / \text{FWHM} = 2.5$ for the SPhP and 7.7 for the waveguide mode, where we used the FWHM of the initial state (a-GST phase) according to literature. These TFOM by far exceed other recently published results in the field of modulated nanophotonics with typical values of 0.5–1.4, and, in the case of the waveguide mode, with an exceptional TFOM of 7.7, even exceed the results on graphene, where values up to 5 have been reported. In the experimental reflectance maps in Figure 3, this switching can be observed as a drastic red-shift of the corresponding dispersion curves. The GST film thickness was chosen such that the frequency shift is maximal for the SPhP. Notably, for the waveguide mode, a slightly larger frequency shift and thus also a larger TFOM is expected for GST films of $\approx 220 \text{ nm}$ thickness (see Figure S1 in the Supporting Information). For even thicker films, the shift decreases, because all modes for both GST phases are pushed against the lower frequency limit given by $\omega_{\text{TO}} = 797 \text{ cm}^{-1}$. For thinner films, on the other hand, the SPhP contrast decreases as well because of a reduced influence of the GST film, while the waveguide mode disappears completely. The film thickness dependence and the boundaries in which the waveguide mode is supported are discussed.
Further below (Figure 4). For the exemplary film thickness of 144 nm, however, our experimental findings clearly demonstrate the tuning potential of both the p-polarized SPhP and the s-polarized waveguide mode supported in GST films on a SiC substrate via GST phase switching.

4. Waveguide Mode Theory

The theoretical description of waveguide modes in a slab waveguide structure is well known and has been given elsewhere. Here, we follow the work by Tien and Ulrich and apply the theoretical model to our air/GST/SiC system, in order to verify the waveguide nature of the s-polarized mode, as well as provide a thorough understanding of the dispersion of this mode and the boundaries in which it is supported. We will mainly focus our discussion on a-GST, which is the GST phase with lower losses, thus featuring better waveguide characteristics. We note, however, that the following theoretical description is equally valid for c-GST. In fact, except for a red-shift of the dispersion and a larger momentum range, the resulting dispersion for c-GST is qualitatively the same as for a-GST (except at the upper momentum boundary, see Figure S2 in the Supporting Information for details). The equation of waveguide modes is given by

$$\frac{\omega}{c_0} \sqrt{\varepsilon_{\text{GST}}} - k^2 d - \Phi_{\text{GST/gst}} = \Phi_{\text{GST/air}} = m \pi$$  \hspace{1cm} (1)

where $\Phi_{\text{GST/gst}}$ and $\Phi_{\text{GST/air}}$ are the phase differences upon reflection at the substrate/film (SiC/GST) and the film/ambient medium (GST/air) interfaces, $\kappa \equiv k/k_0 = n_{\text{GST}} \sin \theta$ is the in-plane momentum (with $k_0 = \omega/c_0$), $d$ is the film thickness, $\omega$ is the frequency, and $c_0$ is the speed of light in vacuum. Please note that Equation (1) is valid for s- and p-polarized waveguide modes. In this work, however, we focus on the zero-order s-polarized waveguide mode and its comparability to the simultaneously supported SPhP mode (for the same frequency, the SPhP is found at larger in-plane momenta). The following discussion will therefore only consider the s-polarized waveguide solutions. The phase differences for s-polarization are given by

$$\tan(\Phi_{\text{GST/gst}}) = \frac{k^2 - \varepsilon_{\text{GST}}}{\sqrt{\varepsilon_{\text{GST}}} - k^2}$$  \hspace{1cm} (2)

$$\tan(\Phi_{\text{GST/air}}) = \frac{k^2 - \varepsilon_{\text{air}}}{\sqrt{\varepsilon_{\text{air}}} - k^2}$$  \hspace{1cm} (3)

In order to ensure total internal reflection inside the waveguide core (here GST), the phase differences $\Phi$ have to be real. As a consequence, the imaginary part of $\Phi$ allows to define those regions in the dispersion where a waveguide mode is supported, and those in which mode propagation is not possible. Figure 4a shows a map of $\Phi_{\text{GST/gst}}$ for the air/a-GST/SiC system. The three white areas (i-iii) are the regions where the a-GST/SiC interface supports the s-polarized waveguide mode ($\text{Im}(\Phi_{\text{GST/gst}}) = 0$). However, for a mode to be supported in the film, the phase differences at both interfaces have to be real simultaneously. Therefore, region (ii) can be ruled out because for in-plane momenta $k > \sqrt{\text{Re}(\varepsilon_{\text{GST}})} = 3.7$ (black dashed line), the phase difference at the a-GST/air interface is complex, $\text{Im}(\Phi_{\text{GST/air}})$ > 0 (see Figure S2 in the Supporting Information). For $k < 3.7$, on the other hand, $\text{Im}(\Phi_{\text{GST/air}}) = 0$ and thus regions (i) and (iii) can support waveguide modes, and their boundaries are solely governed by the a-GST/SiC interface. In fact, the lower boundary along the momentum axis is given by the SiC permittivity $\sqrt{\text{Re}(\varepsilon_{\text{SiC}})}$ (red dashed line), which is a highly dispersive quantity and is the reason for the rich physics of waveguide modes bounded by polar crystal substrates. While in principle, region (iii) does support waveguide modes, we here only focus on region (i) due to its capacity of supporting the s-polarized waveguide mode.
waveguide mode that features an upward bended, polariton-like dispersion in proximity to the SiC substrate SPhP mode (see Figure 1e,f). As for the SPhP, the lower boundary of region (i) along the frequency axis is defined by the TO phonon frequency of SiC, \( \omega_{\text{SiC}} \) (gray dashed line). Interestingly, the reason that \( \omega_{\text{SiC}} \) is a boundary is different for the two modes: The SPhP requires negative \( \text{Re}(\varepsilon_{\text{SiC}}) \) to have an evanescently decaying electric field in z-direction. On the other hand, the waveguide mode requires \( \text{Re}(\varepsilon_{\text{GST}}) < \text{Re}(\varepsilon_{\text{SiC}}) \) to enable total internal reflection and thereby get evanescently decaying fields. For frequencies \( \omega < \omega_{\text{SiC}} \), both conditions are lost simultaneously.

Additional to the restrictions given by total internal reflection that are reflected in the phase differences \( \Phi \) (Figure 4a), waveguide modes in asymmetric slab waveguides require a minimal film thickness \( d_{\text{min}} \) in order to be supported. Evaluation of Equation (1) at the lower and upper momentum boundaries leads to the following expression for \( d_{\text{min}} \) for s-polarized waveguide modes\(^{(37)}\):

\[
d_{\text{min}} = \frac{m\pi + \arctan\left(\frac{\varepsilon_{\text{SiC}} - \varepsilon_{\text{GST}}}{\varepsilon_{\text{GST}} - \varepsilon_{\text{SiC}}}\right)}{\omega} \tag{4}
\]

Note that in asymmetric slab waveguides such as the air/GST/SiC system, the minimal film thickness for the zero-order s-polarized waveguide mode is smaller than for the zero-order p-polarized mode.\(^{(56)}\) As a consequence, for the here studied film thickness range of 0–800 nm, only the s-polarized waveguide mode can be observed.

In Figure 4b, \( d_{\text{min}} \) of the air/a-GST/SiC system is plotted as a function of frequency for the first four orders m. Interestingly, in the SiC reststrahlen band, \( d_{\text{min}} \) of the \( m = 0 \) mode is almost zero, but features a rapid increase at larger frequencies with a maximum value of \( d_{\text{min}} = 226 \text{ nm} \). As a consequence, a-GST films of thicknesses \( d_{\text{GST}} > 226 \text{ nm} \) support the \( m = 0 \) mode at any frequencies, while in thinner films, such as our sample with \( d_{\text{GST}} = 144 \text{ nm} \), the mode is forbidden in a broad frequency range (gray area in Figure 4b). Therefore, the \( m = 0 \) s-polarized waveguide mode we observe in our experiments is limited to a frequency range between 797 and 1056 cm\(^{-1}\), which is very similar to the allowed frequency range of the SPhP defined by the SiC reststrahlen band (797–970 cm\(^{-1}\)).

The range in which the s-polarized air/a-GST/SiC waveguide mode can be observed is best illustrated in the dispersion plot we show in Figure 4c. Here, the gray areas indicate the forbidden regions given by \( \text{Im}(\Phi_{\text{SiC/GST}}) \). For the two thinnest films of 144 and 200 nm (red and orange curves), \( d_{\text{min}} \) defines a frequency range where the \( m = 0 \) mode is forbidden, which is represented by the region in which the two dispersion curves are disrupted. The restriction given by \( d_{\text{min}} \) coincides exactly with the restriction given by \( \text{Im}(\Phi_{\text{SiC/GST}}) \). For the three thicker films (light green, dark green, and blue curves), on the other hand, the mode dispersions are continuous, as there is no minimum film thickness required. However, these mode dispersions are pushed against the lower frequency boundary \( \omega_{\text{SiC}} \) with increasing film thickness, and the dispersion of the mode in the thickest film of 1200 nm even starts to disappear at this boundary. This behavior originates in the pole of the dielectric function of SiC at \( \omega_{\text{SiC}} \), constituting the hard lower barrier of the frequency range in which waveguide modes can be supported. Notably, thanks to this boundary, the waveguide mode dispersion experiences the upward bending that leads to the resemblance of the SPhP mode dispersion. Finally, we note that the shifting of the dispersion with film thickness enables a straightforward way of tuning the mode resonance across the entire allowed frequency range.

5. Surface Polariton-Like Waveguide Modes for s-Polarized Nanophotonics

Equipped with this theoretical understanding, we now turn our focus on the comparability of the SPhP and the thin film s-polarized waveguide mode in terms of optical field enhancement and spatial confinement. Figure 5a–f shows z-profiles of the in-plane electric field enhancement (FE) of the SPhP (\( E_0 \)) and the waveguide mode (\( E_\text{s} \)) at their corresponding resonance frequency for a set of different a-GST film thicknesses, calculated for excitation via a KRS5 coupling prism in the Otto geometry at an incident angle of \( \theta = 30^\circ \). Furthermore, the continuous behavior of the maximum field enhancement \( \text{FE}_{\text{max}} \) for the SPhP (blue curve) as a function of thin a-GST film thickness \( d_{\text{GST}} \) is shown in Figure 5g, and Figure 5h shows the corresponding mode frequency \( \omega_{\text{mode}} \) (\( \text{FE}_{\text{max}} \) and \( \omega_{\text{mode}} \) for larger film thicknesses are shown in Figure S3, Supporting Information).

A bare SiC surface (Figure 5a) only supports an SPhP with a maximum field enhancement (\( \text{FE}_{\text{max}} \)) of \( \approx 7 \), which is, however, reduced when an additional a-GST film of 144 nm is placed on top of SiC (Figure 5b). For increasing film thicknesses (Figure 5 c–e), the field enhancement of the SPhP drops even further and for the largest a-GST film of 800 nm thickness (Figure 5f), there is no enhancement observable at all, meaning that the structure does not support the air/SiC SPhP anymore. This transition toward a total loss of the SPhP for large film thicknesses stems, on the one hand, from the fact that the observed air/SiC SPhP originates in the dielectric contrast of the adjacent air and SiC, and for growing a-GST film thicknesses, there is no SiC/air interface anymore. On the other hand, the TO phonon is IR-active and therefore absorption in SiC increases drastically in the vicinity of \( \omega_{\text{SiC}} \), contributing to the observed loss of field enhancement. Finally, we cannot observe the emergence of any other SPhP for thicker films, because the air/a-GST interface does not feature permittivities of opposite sign, while a potential a-GST/SiC SPhP cannot be excited via prism coupling with a KRS5 prism, because the required momenta are too large (\( n_{\text{KRS5}} < n_{\text{a-GST}} \)).

The s-polarized waveguide mode, on the other hand, features a maximum field enhancement of \( \approx 8 \) in a 250–400 nm thin a-GST film, which decreases for both thinner and thicker films (Figure 5g, red curve). In order to explain this behavior, we recall that for small in-plane momenta (\( 1 < \kappa_\parallel < 2 \)), the waveguide mode is restricted to a frequency range between \( \omega_{\text{SiC}} = 797 \) and 1000–1200 cm\(^{-1}\), where the upper boundary depends on \( \kappa_\parallel \) (Figure 4c). With increasing film thickness, the dispersion red-shifts through the allowed frequency region and eventually is pushed against \( \omega_{\text{SiC}} \), as can be observed in Figure 5h (red curve). Here, analogous to the SPhP, the high absorption in SiC close to \( \omega_{\text{SiC}} \) leads to the observed loss of field enhancement. At the upper frequency boundary, on the other
hand, the permittivity of SiC is close to 1, resulting in a less confined mode in SiC compared to modes at frequencies further into the reststrahlen band (compare the penetration depth into SiC of the waveguide mode in Figure 5b,d), thus leading to smaller field enhancements.

The penetration depth into the SiC substrate \( \delta \), and thus the spatial confinement of the modes, is solely a function of the SiC permittivity tensor. Close to \( \omega_{TO_{SiC}} \), both the SPhP and the waveguide mode are strongly confined (with a penetration depth of \( \delta \approx 100 \text{ nm at } 800 \text{ cm}^{-1} \)), whereas at higher frequencies, the confinement decreases and the modes leak further into the SiC substrate (\( \delta \approx 1.5 \mu\text{m at } 950 \text{ cm}^{-1} \), see Figure S3 in the Supporting Information for details). This frequency dependence of the spatial confinement can also be seen in Figure 5i,j, where we show spatio-spectral maps of the SPhP and the waveguide mode, respectively, showing their different resonance frequencies while featuring comparable Q-factors. The maps are also calculated for an in-plane momentum of \( k/k_0 = 1.18 \). In contrast to (a)–(f), we here plot with a single scale of the z-axis in order to better visualize the evanescent depth of the modes along the z-direction. The z-positions are labeled relative to the respective interface with a-GST.

As we have shown exemplarily for a SiC substrate, a thin overlying GST film enables an s-polarized waveguide mode with features such as its dispersion, field enhancement, and spatial confinement being comparable to those of the simultaneously supported SPhP. Strikingly, this concept can be easily transferred to any polar crystal substrate. In Figure 6, we compare SiC to aluminum nitride (AlN), gallium nitride (GaN), and zinc oxide (ZnO), all featuring a single reststrahlen band, and the multimode material \( \alpha \)-quartz (\( \alpha \)-SiO\(_2\)). For each material, the dispersions of the supported SPhPs (blue and green circles) and s-polarized waveguide modes (red and orange circles) are plotted for a 144 nm thin a-GST and a 132 nm c-GST film, respectively. The vertical gray bars correspond to the FWHM of the mode resonances extracted from calculated spectra of the imaginary part of the corresponding reflection coefficient, than that of the former. The opposite behavior can be observed when the waveguide mode frequency is close to \( \omega_{TO_{SiC}} \) (Figure 5f) and the SPhP mode sits at higher frequencies (for example Figure 5a,b).
the broad IR frequency range of 400–1400 cm$^{-1}$ dispersing above or at the upper edge of the reststrahlen band.

substrate that simultaneously supports a p-polarized SPhP and comprising a subwavelength thin PCM film on a polar crystal.

6. Conclusion

In conclusion, we have presented a generic material system comprising a subwavelength thin PCM film on a polar crystal substrate that simultaneously supports a p-polarized SPhP and an s-polarized waveguide mode. Our experimental data of the exemplary GST/SiC system demonstrate that high-contrast active tuning of both modes can be realized via switching of the GST phase, achieving an exceptional tuning figure of merit of up to 7.7. We have presented a simple theoretical model for the waveguide mode, revealing the similarity of the SPhP and waveguide mode dispersions in the reststrahlen region of the substrate material. Furthermore, by analyzing the electric field distributions across the system interfaces, we have shown that the waveguide mode can feature spatial confinement and field enhancement factors comparable to those of an SPhP, while expanding the allowed frequency range beyond the reststrahlen band. Thus, in addition to the polarization-limited surface polaritons, polariton-like waveguide modes supported by phase-change materials on polar crystal substrates provide a promising complementary building block for actively tunable, low-loss, and omnipolarized nanophotonic applications, such as in-plane metasurfaces or polariton lenses.

7. Experimental Section

Experimental: The 144 nm thick film of the phase-change material Ge$_2$Sb$_2$Te$_5$ was deposited onto the SiC substrate by direct current magnetron sputtering with a background pressure of about 2 $\times$ 10$^{-6}$ mbar and 20 sccm Ar flow in constant power mode (20 W) using stoichiometric targets of 99.99% purity. The experimental setup is illustrated in Figure 2a and has been described before. In short, the Otto geometry was implemented employing three motorized actuators (Newport TRA12ZPD) to control the relative position of the coupling prism (KRS5, 25 mm high, 25 mm wide, angles of 30°, 30°, and 120°, with the 120° edge cut-off, Korth Kristalle GmbH) with respect to the sample. The motors pushed the prism against springs away from the sample and thus enabled continuous tuning of the air gap width $d_{\text{air}}$. The 120° prism edge was cut-off parallel to the backside in order to quantify $d_{\text{cut}}$ via white-light interferometry. The smallest gap width achieved was of $\approx$ 3 µm. In principle, gap widths below 3 µm can be measured, but were not feasible for the GST/SiC sample probably due to a not perfectly flat sample surface.

For each incident angle and polarization, the air gap of critical mode coupling $d_{\text{cut}}$ was determined by optimizing the mode resonance depth in the reflectance signal, and all spectra were taken at $d_{\text{crit}}$. However, $d_{\text{cut}}$ scales with the resonance quality, because large losses into the sample require equally large radiative losses that are provided by the proximate coupling prism, thus leading to very small $d_{\text{cut}}$. This can lead to $d_{\text{cut}}$ being smaller than the experimentally achieved minimum gap of 3 µm, resulting in low resonance depths due to undercoupling. This is, indeed, the reason why the waveguide mode resonance in the lossy c-GST maximally dropped by 30% in the experiments (see Figures 2a and 3), whereas all other resonances achieved 80%.

Furthermore, it is noted that the in-plane electric field distributions of SPhPs and waveguide modes in Figure 1 a–d, analogous to Figure 5a–f, were calculated for excitation via a KR5 coupling prism in the Otto geometry at an incident angle of $\theta$ = 30°. For all four systems, the coupling gap was fixed to the critical coupling gap of the SPhP on bare SiC ($d_{\text{cut}}$ = 5.3 µm). This asymmetric excitation scheme is the reason for the asymmetric field distribution of the air-bound waveguide mode in Figure 1d.

The mid-infrared FEL employed as excitation source was operated at an electron energy of 31 MeV, yielding a tuning range of the FEL output wavelength between 7.7 and 18 µm (1400–550 cm$^{-1}$, covering the spectral range of the SiC reststrahlen band) by moving the motorized undulator gap. The micropulse and micropulse energies were $\approx$ 30 mJ and 10 µJ, respectively.
respectively. However, the incident FEL beam was attenuated by \(-18\ \text{dB}\) in order to avoid irreversible damage of the GST film and unintentional crystallization. Further details on the FEL are given elsewhere.\[40]\n
The reflectance signal was detected with a pyroelectric sensor (Eltec 420M7). A second sensor measured the power reflected off a thin KBr plate placed prior to the setup in the beam. This reference signal was used to normalize all reflectance signals and was measured on a single shot basis in order to minimize the impact of shot-to-shot fluctuations of the FEL. The spectral response function of the setup was determined by taking a reflectance spectrum at large gap widths \(d_{\text{gap}} = 40\ \mu\text{m}\), where the beam was totally reflected at the prism backside across the full spectral range. References were measured for each incident angle and polarization individually, and all reflectance spectra were normalized to their respective response function.

**Global Fitting Procedure**: The spectra in Figure 2b, the dispersion curves in Figures 1e,f, 3, and 6, and the electric field distributions in Figure 5 were all calculated with a \(4 \times 4\) transfer matrix formalism\[43\] that accounts for the anisotropy of the c-cut hexagonal 4H-SiC substrate. For the calculations, most of the material parameters and system variables were set to values from literature or in advance determined values (including the phonon frequencies \(\omega_{\text{TO},\text{o}} = 979\ \text{cm}^{-1}\), \(\omega_{\text{TO},\text{e}} = 788\ \text{cm}^{-1}\), \(\omega_{\text{LO},\text{o}} = 970\ \text{cm}^{-1}\), \(\omega_{\text{LO},\text{e}} = 964\ \text{cm}^{-1}\) where \(o\) stands for ordinary and \(e\) for the extraordinary crystal direction\[35\], the damping \(\delta_{\text{TO}} = 3.75\ \text{cm}^{-1}\), \(\delta_{\text{LO}} = 6.5\ \text{cm}^{-1}\) the spectral width of the FEL \(\Delta \omega = 5.56\ \text{cm}^{-1}\), the angular width of the excitation beam \(\Delta \theta = 0.24^\circ\), and all nine incident angles \(\theta = 26^\circ–34^\circ\). The remaining parameters, on the other hand, were extracted from a global fitting procedure and comprised the a-GST film thickness \(d_{\text{GST}} = 144\ \text{nm}\) (the c-GST film thickness was calculated by a constant factor, \(d_{\text{GST}} = 0.92\ d_{\text{GST}} = 132\ \text{nm}\))\[44\], the real and imaginary part of the permittivity of c-GST \((\epsilon_{\text{c-GST}} = 31.4 + 9)\), the real permittivity of a-GST \((\epsilon_{\text{a-GST}} = 13.8 + 0)\), the gap widths \(d_{\text{gap}}\), and a multiplication factor \(a\) to account for slow FEL power drifts (0.95 < \(a\) < 1). The data sets of all four system configurations (p-polarization and s-polarization for a-GST and c-GST, respectively), each containing nine spectra at different incident angles, were fitted globally, where \(d_{\text{GST}}, \epsilon_{\text{GST}}\) and \(\epsilon_{\text{c-GST}}\) were global fitting parameters, while the factors \(a\) and the gap widths \(d_{\text{gap}}\) were adjusted individually for each spectrum.

**Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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**Conflict of Interest**

The authors declare no conflict of interest.

**Keywords**

active nanophotonics, highly confined waveguide modes, infrared spectral range, phase-change materials, surface phonon polaritons

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[35] T. G. Folland, A. Fali, S. T. White, J. R. Matson, S. Liu, N. A. Aghamiri, J. H. Edgar, R. F. Haglund, Y. Abate, J. D. Caldwell, Nat. Commun. 2018, 9, 4371.

[36] K. Chaudhary, M. Tamagnone, X. Yin, C. M. Spägele, S. L. Oscurato, J. Li, C. Persch, R. Li, N. A. Rubin, L. A. Jauregui, K. Watanabe, T. Taniguchi, P. Kim, M. Wuttig, J. H. Edgar, A. Ambrosio, R. F. Haglund, Y. Abate, J. D. Caldwell, Phys. Rev. B 2018, 98, 201101.

[37] P. K. Tien, R. Ulrich, J. Opt. Soc. Am. 1970, 60, 1325.

[38] A. Paarmann, I. Razdolski, A. Melnikov, S. Gewinner, W. Schöllkopf, M. Wolf, Appl. Phys. Lett. 2015, 107, 081101.

[39] N. C. Passler, I. Razdolski, S. Gewinner, W. Schöllkopf, M. Wolf, A. Paarmann, ACS Photonics 2017, 4, 1048.

[40] W. Schöllkopf, S. Gewinner, H. Junkes, A. Paarmann, G. von Helden, H. Bluem, A. M. M. Todd, Proc. SPIE 2015, 9512, 95121L.

[41] W. K. Njoroge, H.-w. Wöltgens, M. Wuttig, J. Vac. Sci. Technol. A 2002, 20, 230.

[42] A.-K. U. Michel, Ph.D. Thesis, RWTH Aachen University 2016.

[43] N. C. Passler, A. Paarmann, J. Opt. Soc. Am. B 2017, 34, 2128.

[44] T. Wang, P. Li, B. Hauer, D. N. Chigrin, T. Taubner, Nano Lett. 2013, 13, 5051.

[45] J. D. Caldwell, O. J. Glembocki, Y. Francescato, N. Sharac, V. Giannini, F. J. Bezarest, J. P. Long, J. C. Owrutsky, I. Vurgaftman, J. G. Tischler, D. S. Kates, K. Watanabe, T. Taniguchi, P. Kim, M. Wuttig, J. H. Edgar, A. Ambrosio, R. F. Haglund, Y. Abate, J. D. Caldwell, Nano Lett. 2018, 18, 4285.

[46] H. Sumikura, T. Wang, P. Li, A.-K. U. Michel, A. Heßler, L. Jung, M. Lewin, M. Wuttig, D. N. Chigrin, T. Taubner, Nano Lett. 2019, 19, 2549.

[47] J. D. Caldwell, A. V. Kretinin, Y. Chen, V. Giannini, M. M. Fogler, Y. Francescato, C. T. Ellis, J. G. Tischler, C. R. Woods, A. J. Giles, M. Hong, K. Watanabe, T. Taniguchi, S. A. Maier, K. S. Novoselov, Nat. Commun. 2014, 5, 5221.

[48] Y. Chen, Y. Francescato, J. D. Caldwell, V. Giannini, T. W. W. Maß, O. J. Glembocki, F. J. Bezarest, T. Taubner, R. Kasica, M. Hong, S. A. Maier, ACS Photonics 2014, 1, 718.

[49] M. Tamagnone, K. Chaudhary, A. Zhu, M. Meretska, J. Li, J. H. Edgar, A. Ambrosio, F. Capasso, 2019, arXiv:1905.02177.

[50] Y. Yao, M. A. Kats, P. Genevet, N. Yu, Y. Song, J. Kong, F. Capasso, Nano Lett. 2013, 13, 1257.

[51] G. Si, E. S. P. Leong, X. Jiang, J. Lv, J. Lin, H. Dai, Y. J. Liu, Phys. Chem. Chem. Phys. 2015, 17, 13223.

[52] S. Dai, J. Zhang, Q. Ma, S. Kittiwatanakul, A. McLeod, X. Chen, S. G. Corder, K. Watanabe, T. Taniguchi, J. Lu, Q. Dai, P. Jarillo-Herrero, M. Liu, D. N. Basov, Adv. Mater. 2019, 31, 1900251.

[53] Z. Fang, S. Thongrattanasiri, A. Schlather, Z. Liu, L. Ma, Y. Wang, P. M. Ajayan, P. Nordlander, N. J. Halas, F. J. Garcia de Abajo, ACS Nano 2013, 7, 2388.

[54] W. Gao, G. Shi, Z. Jin, J. Shu, Q. Zhang, R. Vajtai, P. M. Ajayan, J. Kono, Q. Xu, Nano Lett. 2013, 13, 3698.

[55] F. Engelbrecht, R. Helbig, Phys. Rev. B 1993, 48, 15698.

[56] S. O. Kasap, Optoelectronics & Photonics: Principles & Practices, 2nd ed., Pearson, London 2013, p. 544.

[57] H. Kogelnik, in Integrated Optics (Ed. T. Tamir), Springer, Berlin 1975. Ch. 2, pp. 13–81.

[58] N. C. Passler, C. R. Gubbin, T. G. Folland, I. Razdolski, D. S. Katzer, D. F. Storm, M. Wolf, S. De Liberato, J. D. Caldwell, A. Paarmann, Appl. Phys. Lett. 2018, 113, 161103.

[59] B. Neuner III, D. Korobkin, C. Fietz, D. Carole, G. Ferro, G. Shvets, Opt. Lett. 2009, 34, 2667.