Infrared Permittivity of the Biaxial van der Waals Semiconductor $\alpha$-MoO$_3$ from Near- and Far-Field Correlative Studies

Gonzalo Álvarez-Pérez, Thomas G. Folland, Ion Errea, Javier Taboada-Gutiérrez, Jiahua Duan, Javier Martín-Sánchez, Ana I. F. Tresguerres-Mata, Joseph R. Matson, Andrei Bylinkin, Mingze He, Weiliang Ma, Qiaoliang Bao, José Ignacio Martín, Joshua D. Caldwell,* Alexey Y. Nikitin,* and Pablo Alonso-González*

The biaxial van der Waals semiconductor $\alpha$-phase molybdenum trioxide ($\alpha$-MoO$_3$) has recently received significant attention due to its ability to support highly anisotropic phonon polaritons (PhPs)—infrared (IR) light coupled to lattice vibrations—offering an unprecedented platform for controlling the flow of energy at the nanoscale. However, to fully exploit the extraordinary IR response of this material, an accurate dielectric function is required. Here, the accurate IR dielectric function of $\alpha$-MoO$_3$ is reported by modeling far-field polarized IR reflectance spectra acquired on a single thick flake of this material. Unique to this work, the far-field model is refined by contrasting the experimental dispersion and damping of PhPs, revealed by polariton interferometry using scattering-type scanning near-field optical microscopy (s-SNOM) on thin flakes of $\alpha$-MoO$_3$, with analytical and transfer-matrix calculations, as well as full-wave simulations. Through these correlative efforts, exceptional quantitative agreement is attained to both far- and near-field properties for multiple flakes, thus providing strong verification of the accuracy of this model, while offering a novel approach to extracting dielectric functions of nanomaterials. In addition, by employing density functional theory (DFT), insights into the various vibrational states dictating the dielectric function model and the intriguing optical properties of $\alpha$-MoO$_3$ are provided.
predictive capabilities for advanced optical devices, it is imperative to develop an accurate dielectric function model. As such, prior work using α-MoO3 has relied on a dielectric function that was estimated from previously reported optic phonon frequencies.\(^{[10,20,21]}\) This approach i) assumes the accuracy of these prior reports of the phonon frequencies, and ii) cannot accurately extract the high- and low-frequency dielectric constants nor the phonon damping constants. These parameters are critical not only for building an accurate model, but for providing insights into how the crystal lattice dictates the optical response of the material. Here, we use model fitting of polarized far-field IR spectroscopy on single thick flakes of α-MoO3 to overcome these limitations, identifying the phonon frequencies, high- and low-frequency dielectric constants, and damping rates. Moreover, in an effort to refine this far-field approach and to compensate for any errors induced by the limited lateral scale of the flake (150 × 650 µm), which prevents conventional ellipsometry measurements, we compare near-field measurements of the damping and the dispersion of PhPs propagating on a thin flake of α-MoO3 to full-wave numerical simulations, analytical dispersions and transfer-matrix calculations that use the initially extracted dielectric function as input. Exceptional quantitative agreement is simultaneously attained between the model and far- and near-field results through an iterative process combining both correlated efforts, dramatically improving the fit and thereby strongly verifying the accuracy and robustness of the resulting dielectric function. Additionally, we employ DFT to give insights into the vibrational modes dictating the oscillators on which this dielectric function model is based and into the intriguing optical properties of the material. Our work therefore reports an accurate and predictive model for the IR dielectric function of α-MoO3, an emerging vdW material for nanophotonics,\(^{[22,23]}\) while also offering an innovative approach to extracting dielectric functions of nanomaterials, where the use of traditional methods is challenging\(^{[24–26]}\) or even not possible.

The exotic properties of PhPs in α-MoO3 stem from their strong anisotropy, a consequence of the crystal structure, which is orthorhombic (Figure 1a). In it, layers formed by distorted α-MoO6 octahedra are weakly bound by vdW forces and all three lattice constants \((a = 0.396 \text{ nm, } b = 1.385 \text{ nm, and } c = 0.369 \text{ nm})\) of α-MoO3 crystals typically appear to be rectangular owing to the anisotropic crystal structure. Labeled arrows indicate crystal directions.

\[
\alpha-\text{MoO}_3\text{. a. Sketch of the unit cell of } \alpha-\text{MoO}_3\text{ and correspondence between the crystallographic directions [100], [001], }\alpha\text{[010], and the spatial coordinates } x, y, z; \text{ the lattice constants are } a = 0.396 \text{ nm, } b = 1.385 \text{ nm, and } c = 0.369 \text{ nm. The blue/red spheres represent molybdenum/oxygen atoms. b) Optical image of an } \alpha-\text{MoO}_3\text{ flake on top of AgCl. } \alpha-\text{MoO}_3\text{ crystals typically appear to be rectangular owing to the anisotropic crystal structure. Labeled arrows indicate crystal directions.}
\]

Traditionally, the dielectric function of a material is extracted using far-field polarized reflection and transmission or ellipsometry measurements. In this work, to extract the three principal values \(\varepsilon_r(\omega), \varepsilon_i(\omega), \text{ and } \varepsilon_z(\omega)\) of α-MoO3 in the mid- and far-IR —between \(\sim9\) and \(\sim23 \mu\text{m}\) free-space wavelengths (1111–435 cm\(^{-1}\) )—we initially carry out Fourier transform infrared (FTIR) micro-spectroscopy. This technique can capture the optical response of relatively small single crystals across a broad spectral bandwidth, as required for accurate and unique determination of the dielectric function of α-MoO3. The process for using FTIR micro-spectroscopy to determine the dielectric function involves taking accurate reflection and transmission spectra from a flake, and then using a Kramers–Kronig consistent model to fit to the experimental results. This model is typically chosen as it provides useful insights into the vibrational and/or electronic states of the material.

While in principle FTIR micro-spectroscopy can determine the optical properties of extremely small—down to 10 µm × 10 µm—flakes, in practice this limits the spectral range, accuracy, and signal-to-noise ratio of spectra, in comparison to measurements on large films, due to the diffraction limit of light. As α-MoO3 is a biaxial crystal with optic
phonons that occur in the LWIR ($\lambda > 9.9$ $\mu$m), we acquired polarized FTIR reflection (Figure 2) and transmission (Supporting Information) spectra from the same relatively large area of an $\alpha$-MoO$_3$ single flake (red square in Figure 1b), to ensure that good quality data were obtained across the full spectral range of interest, with the polarizer aligned along both in-plane principal axes (see Experimental Section for the detailed experimental procedure). The flake, with lateral dimensions of about 150 $\mu$m by 650 $\mu$m and a thickness of approximately 3 $\mu$m (Supporting Information), was exfoliated from bulk $\alpha$-MoO$_3$ crystals and then transferred onto an AgCl substrate (Figure 1b), which is highly transparent across the
entire spectral range of interest (see dielectric function in the Supporting Information).

As a result of the strong in-plane anisotropy in the optical response of $\alpha$-MoO$_3$, the reflectance spectra along the two in-plane directions, [001] and [001], are clearly different (Figure 2). There are several high-reflectivity bands covering different frequency ranges, termed RBs, which are originated by the polar nature of the bands in the so-called polar crystals, which include $\alpha$-MoO$_3$. These bands result from the breaking of the degeneracy between the longitudinal optical (LO) and transverse optical (TO) phonons, and the latter becoming IR active. Consequently, a strong absorption occurs at the TO phonon, and the real part of the frequency-dependent dielectric permittivity becomes negative within the RB, leading to the well-reported high reflectivity of polar materials within this spectral region.[28] Thus, we can use the reflectance data to approximate the position of the TO and LO phonons by the onset of the RB and one oscillator for $\varepsilon_x$ and $\varepsilon_z$, respectively:

$$
\varepsilon_x(\omega) = \varepsilon_x^\infty + \frac{\varepsilon_x^L - \varepsilon_x^T}{(\omega^L - i\gamma_x^L) - (\omega^T - i\gamma_x^T)},
$$

$$
\varepsilon_z(\omega) = \varepsilon_z^\infty + \frac{\varepsilon_z^L - \varepsilon_z^T}{(\omega^L - i\gamma_z^L) - (\omega^T - i\gamma_z^T)},
$$

where $\varepsilon_i(\omega)$ denotes the $i$-th principal component of the permittivity tensor, and $i = x, y, z$ stand for the three principal axes of the crystal, which correspond to the crystalline directions of $\alpha$-MoO$_3$, [001], [001], and [010], respectively. $\gamma_i$ represents the high-frequency dielectric constant, while $\omega_i^L$ and $\omega_i^T$ refer to the LO and TO phonon frequencies, along the $i$-th direction. $\gamma_i$ represents the damping factor of the Lorentzian line shape that is derived from the phonon scattering rate and is inversely proportional to the phonon scattering lifetime. Finally, $j$ is the subscript denoting the different phonon pairs along the same axis.

Using this model and our experimental data, we can in principle extract all the parameters that define the IR dielectric function in Equation (2) (Experimental Section), as previously employed for measuring the dielectric function of other anisotropic vdW materials, as hBN[28,31] or quartz.[32,33] However, this approach relies heavily upon reflection spectra collected at low angles, which is not very sensitive to absorption and thus damping, making the fitting procedure weakly sensitive to the phonon damping $\gamma_i$. Furthermore, experimental non-idealities influence the fitted $\varepsilon_x^T$ and LO phonon energy values (Supporting Information). On the other hand, near-field polariton imaging by s-SNOM is extremely sensitive to both damping and LO phonon energies. More specifically, near the LO phonon frequency, the PhP wavelength and propagation length from s-SNOM measurements on thin flakes at multiple incident laser frequencies, these parameters can be determined with dramatically improved accuracy. Even so, s-SNOM is not highly sensitive to other parameters, such as the high-frequency permittivity, and it is difficult to obtain measurements close to the TO phonon due to strong absorptive losses. Therefore, our procedure for extracting the permittivity of $\alpha$-MoO$_3$ is based on an iterative effort utilizing the two correlated measurements in the far- and near-field, respectively.

Particularly, the free parameters, $\varepsilon_x^T$, $\omega_x^L$, $\omega_x^T$, and $\gamma_x$, in Equation (2), are initially adjusted so that the simulated reflectance more accurately reproduces the experimental far-field FTIR spectra (Figure 2). To do so, the system is modeled as a three-layer structure (air/$\alpha$-MoO$_3$/AgCl). After this first adjustment of the parameters (Supporting Information), they are provided as inputs to model near-field experimental data. For this purpose, we use the recently derived analytical dispersion of polaritons propagating in a biaxial slab embedded between two semi-infinite media[34]

$$
q = \frac{\rho}{k_d} \left[ \arctan \left( \frac{\varepsilon_1 \rho}{\varepsilon_x} \right) + \arctan \left( \frac{\varepsilon_1 \rho}{\varepsilon_x} \right) + \pi \right], l = 0, 1, 2...
$$

where $q = k_i/k_0$ is the normalized in-plane momentum ($k_i = k^L + k^T$), $\varepsilon_1$ and $\varepsilon_x$ are the permittivities of the superstrate and substrate, respectively, $d$ is the thickness of the $\alpha$-MoO$_3$ flake, $k_0 = \omega/c$ is the wavevector in free space and $\rho = i\varepsilon_1 \varepsilon_x / (\varepsilon_1 \cos \alpha + \varepsilon_x \sin^2 \alpha)$, with $\alpha$ being the angle between the $x$ axis and the in-plane component vector. On the other hand, an instructive way to visualize both the dispersion and the damping is via a false-color plot of the imaginary part of the Fresnel reflection coefficient, $\text{Im}(r_0)$, at real $q$ and $\omega$, obtained by means of transfer-matrix calculations.[35] Briefly, PhPs correspond to the divergences of the calculated reflectivity $r_0(q, \omega)$ of the anisotropic structure at complex momenta $q$. Both the analytical dispersion from Equation (3) and that inferred from transfer-matrix calculations are compared to the experimental dispersion (Figure 3), measured via polariton interferometry on an $\alpha$-MoO$_3$ flake of thickness $d = 120$ nm (Supporting Information) placed on top of BaF$_2$, which, as AgCl, is highly transparent across the entire spectral range of interest (see dielectric function in Supporting Information), allowing for a finer tuning of the free parameters in Equation (2). Following this approach, an excellent agreement between the experiment and our model is obtained. Remarkably, the ability to successfully fit the polaritonic response of a thin $\alpha$-MoO$_3$ flake placed on a different substrate reinforces our claims of robust quantitative
modeling of the far- and near-field IR response of this material, especially considering the strong thickness dependence of hyperbolic polaritons.[17]

To accurately adjust the damping and unambiguously verify that our dielectric function successfully accounts for the polaritonic response of α-MoO3, we analyze the PhP propagation length from both simulated and experimental near-field images. To do so, we use our dielectric function model as input for full-wave numerical simulations (Figure 4b) that mimic s-SNOM measurements (Figure 4a) (see Experimental Section), carried out on a 120-nm-thick α-MoO3 flake on top of BaF2 at two representative frequencies: i) \( \varepsilon_b = 910 \) cm\(^{-1}\)—within the lowest-frequency high-reflectivity band along the [100] crystal direction in Figure 2e (hyperbolic band[9,10] RB\(_2\), 821–963 cm\(^{-1}\)); see Figure 4c,d—and ii) \( \varepsilon_b = 990 \) cm\(^{-1}\)—within the high-reflectivity band along both in-plane directions in Figure 2e (elliptic band[9,10] RB\(_1\), 957–1007 cm\(^{-1}\)); see Figure 4e.f. Within RB\(_1\), we experimentally observe (Figure 4c) fringes along the [100] crystal direction, which indicate the excitation of in-plane hyperbolic PhPs in α-MoO3, consistently with what has been previously reported.[9,10] By extracting a line profile from Figure 4c and fitting it to an exponentially-decaying oscillating function corrected by geometrical spreading factors (Supporting Information), we obtain a PhP wavelength of \( \lambda_p^x = 820 \pm 25 \) nm and a propagation length of \( L_p^x = 1400 \pm 100 \) nm. To account for these experimental results with our dielectric function, we adjust the parameter \( \gamma_{32} \) in the model—which determines the damping of PhPs in RB\(_2\)—and run a full-wave numerical simulation using the adjusted dielectric function. From the image resulting from this simulation (Figure 4d), we find the PhP wavelengths and propagation lengths to be \( \lambda_p^x = 325 \pm 25 \) nm and \( L_p^x = 800 \pm 100 \) nm along the [100] crystal direction and \( \lambda_p^y = 400 \pm 25 \) nm and \( L_p^y = 1100 \pm 100 \) nm along [001] direction. Again, we obtain excellent agreement with the experiment, unambiguously demonstrating the validity of our model.

As a further verification, we also carried out attenuated total reflectance (ATR) spectroscopy of an α-MoO3 flake on top of BaF2 (Supporting Information). FTIR-ATR has been proven to be an effective method for studying the properties of anisotropic 2D materials,[36] as it is sensitive to both dielectric and polaritonic resonant modes supported by these crystals. The good agreement (Supporting Information) between calculated and simulated spectra (using our dielectric function) again demonstrates the validity of our model and its broad applicability for predictive simulations.

We show in Table 1 the resulting frequencies and damping rates of the phonons of our model along all three principal axes and plot the extracted dielectric function in Figure 5 (middle panel). The parameters provided in Table 1 are closely related with prior experimental results[7–9] and previously reported phonon frequencies estimations[20,21] (Supporting Information). However, unlike earlier results, we identified two additional weak phonon bands aligned along the x-axis (100) crystal direction, and accurately determined the high-frequency dielectric permittivity along both the [100] and [001] directions. We note that the weak phonon spectrally located at 998 cm\(^{-1}\) results in a discontinuity in the hyperbolic polariton dispersion relation (Figure 3), which has not been predicted using earlier experimental or theoretical models. We also note that the apparent strong damping of the optic phonon located near 506 cm\(^{-1}\) is likely an artifact resulting from being spectrally near the cutoff of the FTIR mercury cadmium telluride (MCT) detector. As expected for a biaxial crystal, α-MoO3 exhibits different permittivities along the three principal axes throughout the entire spectral range studied (\( \varepsilon_x \neq \varepsilon_y \neq \varepsilon_z \)). Furthermore, over the spectral range extending from 545 to 1006 cm\(^{-1}\), there is always at least one component i exhibiting a negative Re(\( \varepsilon_i \)) value, thus...
the crystal can support PhPs over this entire frequency range, having multiple hyperbolic regimes, each of them when at least one component $i$ fulfills $\text{Re}(\epsilon_i) < 0$.

To identify the vibrational character of the multiple optic phonons supported by $\alpha$-MoO$_3$, and to further test the validity of our dielectric function model, we have also calculated the

Figure 4. Near-field simulated and experimental images of in-plane anisotropic PhPs on an $\alpha$-MoO$_3$ flake. a) Schematics of the s-SNOM experimental configuration and b) the full-wave numerical simulation model mimicking the experiment. c) Experimental near-field amplitude $s_2$ images and d) simulated $z$ component of the electric field $|E_z|$ of a 120-nm-thick $\alpha$-MoO$_3$ flake on top of BaF$_2$ at incident frequency $\omega_0 = 910$ cm$^{-1}$. e,f) Corresponding images for incident frequency $\omega_0 = 990$ cm$^{-1}$.
permittivity tensor and vibrational eigenmodes using DFT (Supporting Information). The resulting ab initio permittivity function qualitatively agrees with the experimentally extracted dielectric function (Figure 5, lower panel). This underlines the potential of employing first-principles methods to characterize the IR permittivity of emerging materials for nanophotonics. However, slight spectral shifts in the phonon frequencies are observed in the calculations with respect to the experiment, which is typical for DFT with semi-local exchange-correlation functionals for semiconducting oxides. A further discrepancy is that, in the ab initio-calculated dielectric function, RB2 and RB3 (Figure 5) do not spectrally overlap due to small underestimation and overestimation of the phonon frequencies along the [100] and [010] directions, respectively, in contrast with the experimental results. We highlight that the ab initio model also predicts several phonon modes beyond the spectral range of our experimental measurements, extending into the THz (Supporting Information). The validity of these predicted modes was verified through experimentally measured far-IR transmission spectra collected from α-MoO3 powder using a Nb-superconducting bolometer (Supporting Information). While we are not able to determine the exact frequencies for the phonons in this spectral band, the combination of the ab initio model and THz transmission spectra suggest that this might be an interesting regime for future work.

In summary, we have extracted the full IR complex dielectric function of the biaxial vdW semiconductor α-MoO3 along all the three crystallographic directions. Its robustness is demonstrated by successfully numerically reproducing three different experimental measurements of flakes with significantly different thicknesses on different substrates. As the index of refraction of the substrate has been recently demonstrated to play a significant role influencing hyperbolic polariton propagation, this further demonstrates the general nature of our reported dielectric function model. We achieved this by combining the virtues of far-field (broadband characterization and accurate determination of TO phonon energies) with near-field imaging (sensitivity to phonon damping and LO phonon energies) and advanced theoretical and numerical approaches, providing an alternative, more accurate approach to predicting the IR dielectric function—and thus the optical response—of 2D, nano and low-dimensional materials. Using the extracted permittivity, we envision that future experiments on IR nano- and optoelectronic devices based on α-MoO3 will be streamlined and optimized using this more accurate model. Based on the breadth of exotic phenomena this material exhibits, such as the first reports of in-plane hyperbolicity within a natural, low-loss crystal and perspectives on negative refraction or hyperlensing and hyperfocusing, the potential for unprecedented planar nanophotonic technologies is envisioned. Building on prior efforts utilizing local changes in the dielectric environment, advanced concepts for reconfigurable planar metaoptics can be realized. More broadly, structuring biaxial materials may offer new opportunities for making both passive and light emitting structures with unusual polarization states—as in the creation of circularly polarized light. Furthermore, it may open new regimes for highly anisotropic dielectric resonators within the highly dispersive, extreme permittivity regime at frequencies below the TO phonon.

| Main axis | Mode index | Correlative far- and near-field experiments | Ab initio calculation |
|-----------|------------|-------------------------------------------|----------------------|
|           |            | $\omega_i$ TO [cm$^{-1}$] | $\omega_i$ LO [cm$^{-1}$] | $\gamma_i$ [cm$^{-1}$] | $\omega_i$ TO [cm$^{-1}$] | $\omega_i$ LO [cm$^{-1}$] | $\gamma_i$ [cm$^{-1}$] |
| x         | 1          | 506.7 | 534.3 | 49.1 | 449 | 467 | 8.3 |
| x         | 2          | 821.4 | 963.0 | 6.0 | 769 | 947 | 3.7 |
| x         | 3          | 998.7 | 999.2 | 0.35 | 1016 | 1018 | 0.4 |
| y         | 1          | 544.6 | 850.1 | 9.5 | 505 | 820 | 12 |
| z         | –          | –    | –     | –   | 765 | 772 | 3.7 |
| z         | 1          | 956.7 | 1006.9 | 1.5 | 976 | 1027 | 0.4 |

“–” : not considered.

Table 1. Parameters for the IR dielectric function of α-MoO3, extracted from correlative far- and near-field experiments (left) and ab initio calculations (right). The high-frequency permittivities are $\varepsilon_{\infty}^x = 5.78$, $\varepsilon_{\infty}^y = 6.07$, and $\varepsilon_{\infty}^z = 4.47$ (fixed to the ab initio value) for the experiment and $\varepsilon_{\infty}^x = 5.86$, $\varepsilon_{\infty}^y = 6.59$, $\varepsilon_{\infty}^z = 4.47$ for the ab initio calculation. The flake thickness used in the fit was 2.9 µm.

Experimental Section

Fourier-Transform Infrared Spectroscopy: Mid-infrared reflectance measurements were undertaken using a Bruker Hyperion 2000 microscope coupled to a Bruker Vertex70v FTIR spectrometer, equipped with a broadband MCT detector (400–8000 cm$^{-1}$), and a wide range FIR beam-splitter (30–6000 cm$^{-1}$). Off-normal ($\times$ 36 Cassegrain, 25° average incidence angle) polarized reflection and transmission incidence spectra were obtained from the crystals. Both KRS5 and polyethylene wire grid polarizers were used in order to optimize the spectral throughput of the system at the relevant phonon frequencies. The spectra were collected with a 2 cm$^{-1}$ spectral resolution and spatial resolution defined by the internal adjustable aperture of the microscope that was set to the size of the specific crystal of interest. All measurements were performed in reference to a gold film.

For FTIR-ATR micro-spectroscopy a modified Bruker optics ATR objective was used. In brief, the objective was modified to reduce its incident angle, and azimuthal angular spread allowing to produce s- and p-polarized light at approximately 39° incidence angle. The Ge (100 µm diameter) prism was pressed into the substrate with a force of approximately 1N, allowing to collect ATR spectra referenced to air. Spectra were taken with the crystal in two orientations with respect to the prism, allowing us to probe both crystal axes, under both polarization states of light.

In addition to the measurements on the α-MoO3 crystals, the dielectric functions of AgCl and BaF$_2$ substrates were determined using unpolarized FTIR spectroscopy. FTIR reflectance and transmission spectra were collected at an angle of 40° using a linear DLaTGs detector, under vacuum conditions. A broader description of such methods as they pertain to polaritonic materials can be found in a recent tutorial.
Fitting of Far-Field Experimental Data: All dielectric function extraction was performed using the WVASE software (J.A. Woolam). Details on the obtained dielectric function are presented in the main text of the manuscript. Unlike infrared ellipsometry, there are a series of non-idealities which need to be accounted for when extracting the dielectric function from FTIR microscopy data. First, we account for the incident angle and spread of the Cassegrain objective, (25° weighted average, 10° spread). Second, light from a Cassegrain objective is distributed over the full 180° azimuthal incidence spread, which means that s- and p-polarized light must be mixed during the fitting process. Finally, due to non-uniformity of the crystal surface, not all incident light was collected, so a spectrally uniform linear scaling factor of 0.92 was used to change the percentage of collected light. Each of these three considerations was required to get an accurate fit of the model. Whilst ideally both reflection and transmission data would be considered for the fit, chromatic aberration induced by the dispersion of the substrate prevented the use of transmission data in our fit. Starting values for the fit were based off those from the ab initio model, and the fit was performed accounting for results along both crystal axes at the same time. The fit was conducted by varying each parameter of the fit systematically, and once sufficiently close a global fit was performed to obtain the fully optimized parameters.

**Scattering-Type Scanning Near-Field Optical Microscopy (s-SNOM):** Infrared nano-imaging was performed using a commercially available s-SNOM (from Neaspec) where a metallized cantilevered atomic force microscope (AFM) tip was used as a scattering near-field probe. The

![Figure 5. IR dielectric function of α-MoO₃. Experimental (middle panel) and ab initio (lower panel) permittivity of α-MoO₃ along the three crystallographic axes. The shaded regions indicate the three Reststrahlen bands (RB₁, RB₂, and RB₃), corresponding to phonons on each of the crystallographic axes: [001], [100], and [010]. The lattice vibrations of these phonons are illustrated in the upper panel (molybdenum/oxygen atoms in blue/red, respectively).](image-url)
tip oscillates vertically at the mechanical resonant frequency (around 270 kHz) of the cantilever, with an amplitude of about 50 nm. The tip is side-illuminated with p-polarized infrared light of frequency $\omega_0$ (from tunable CO2 and quantum cascade lasers) and electric field $E_0$. Acting as an infrared antenna, the Pt-coated tip concentrates the incident field into a nanoscale spot at the apex, which interacts with the sample surface and thus modifies the tip-scattered field $E_{scat}$. $E_{scat}$ is recorded with a pseudo-heterodyne Michelson interferometer. Demodulation of the interferometric detector signal at the n-th harmonics of the tip oscillation frequency yields the complex-valued near-field signals $\sigma_n = \sigma_n e^{i\phi_n}$, with $\sigma_n$ being the near-field amplitude and $\phi_n$ being the near-field phase. By recording the near-field signals as a function of the lateral tip position, we obtain near-field images or line trace. In the case of probing a material supporting polaritons, the nanoscale “hotspot” at the tip apex acts as a local source of polaritons. The tip-launched polaritons reflect at the flake edges and produce polariton interference, yielding fringes in the near-field images. The distance between the interference fringes corresponds to half the polariton wavelength, $\lambda_{pol}/2$.

Full-Wave Numerical Simulations: In s-SNOM experiments the tip acts as an optical antenna that converts the incident light into a strongly confined near field below the tip apex, providing the necessary momentum to excite PhPs. However, owing to the complex near-field interaction between the tip and the sample, numerical quantitative studies of s-SNOM experiments meet substantial difficulties in simulating near-field images. To overcome these difficulties, we approximate the tip by a dipole source (with a constant dipole moment), in contrast to the usual dipole model, in which the effective dipole moment is given by the product of the exciting electric field and the polarizability of a sphere. We assume that the polarizability of the dipole is weakly affected by the PhPs excited in the $\alpha$-MoO3 flake, and their back-action onto the tip can be thus neglected. Calculating the amplitude of the near field, $|E_n|$, as a function of the dipole position $(x,y)$, we simulate near-field images (we use COMSOL MULTIPHYSICS). The experimental s-SNOM images are well reproduced by our simulated images (see Figure 4c–f), which lets us conclude that the calculated field between the dipole and the $\alpha$-MoO3 flake, $E_n$, provides a valid numerical description of the signals measured by s-SNOM.

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

Acknowledgements
G.-P. and J.-T.C. acknowledge support through the Severo Ochoa Program from the Government of the Principality of Asturias (grants No. PA-20-PF-BP19-053 and PA-18-PF-BP17-126, respectively). I.E. acknowledges support from the Spanish Ministry of Economy and Competitiveness (FIS2016-76617-P). J.M.-S. acknowledges support through a Clarín Marie Curie-COFUND grant from the Government of the Principality of Asturias and the EU (PA-18-ACB17-29), and the Ramón y Cajal Program (RYC2018-026196-I) from the Government of Spain. Q.B. acknowledges support from Australian Research Council (ARC, FT150100440, 1H150100006, and CE170100039). J.D.C. was supported by the National Science Foundation (U.S.A.) under grant number U0048926. A.Y.N. acknowledges the Spanish Ministry of Science, Innovation and Universities (national project MAT2017-88358-C3-3-R) and Basque Government (grant No. IT1164-19). PA-G. acknowledges support from the European Research Council under starting Grant 715496, 2DNANOPTICA.

Conflict of Interest
The authors declare no conflict of interest.

Keywords
dielectric function, hyperbolic phonon polaritons, van der Waals materials

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