Long-lived phonon polaritons in hyperbolic materials

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Natural hyperbolic materials with dielectric permittivities of opposite sign along different principal axes can confine long-wavelength electromagnetic waves down to the nanoscale, well below the diffraction limit. This has been demonstrated using hyperbolic phonon polaritons (HPP) in hexagonal boron nitride (hBN) and α-MoO₃, among other materials. However, HPP dissipation at ambient conditions is substantial and its fundamental limits remain unexplored1,2. Here, we exploit cryogenic nano-infrared imaging to investigate propagating HPP in isotopically pure hBN and naturally abundant α-MoO₃ crystals. Close to liquid-nitrogen temperatures, the losses for HPP in isotopic hBN drop significantly, resulting in propagation lengths in excess of 25 micrometers, with lifetimes exceeding 5 picoseconds, thereby surpassing prior reports on such highly-confined polaritonic modes. Our nanoscale, temperature-dependent imaging reveals the relevance of acoustic phonons in hyperbolic polariton damping and will be instrumental in mitigating such losses for miniaturized middle infrared technologies operating at the liquid-nitrogen temperatures.
We investigate hyperbolic phonon polariton (HPP) propagation and dissipation in isotopically pure hexagonal boron nitride (h\textsuperscript{10}BN, h\textsuperscript{11}BN), along with naturally abundant hBN and in α-MoO\textsubscript{3} van der Waals crystals using near-field infrared (IR) microscopy. Nano-IR methods allow one to directly visualize polaritonic standing waves on the surface of these hyperbolic materials\textsuperscript{1-6}. Similar to graphene plasmon polariton investigations\textsuperscript{7-12}, both the wavelength of the HPP \( \lambda_p(\omega) \) and its dissipation \( \gamma \) can be readily obtained from nano-IR images (Figs.1-2), with these two quantities allowing one to extract the complex dielectric function \( \varepsilon(\omega) = \varepsilon' + i \varepsilon'' \) of a phononic medium\textsuperscript{3-6}. Earlier attempts have characterized HPP in isotopically pure hBN (h\textsuperscript{10}BN, h\textsuperscript{11}BN) and in α-MoO\textsubscript{3} crystals at ambient conditions\textsuperscript{6,13-18}. However, the fundamental limits of HPP dissipation and lifetime remain to be determined as this necessarily relies on temperature-dependent nano-imaging of polaritonic waves. In this work, we report cryogenic nano-imaging results for these van der Waals materials for the first time, demonstrating record-long HPP lifetimes. We studied isotopically pure hBN and biaxial hyperbolic α-MoO\textsubscript{3} crystals, and compares these results with naturally abundant hBN crystals. Combined with theoretical models, we examined the physics governing HPP dissipation in isotopically pure hyperbolic crystals.

To perform cryogenic nano-imaging, we utilized a home-built scanning near-field IR microscope operating at variable temperatures\textsuperscript{7,19-21}. In this setup, the incident light with IR frequency \( \omega \) is focused onto the metallized tip of an atomic force microscope (AFM). As the tip approaches the sample, a concentrated evanescent field excites polaritonic modes with a \( \lambda_p(\omega) \) that is much shorter than the free-space wavelength \( \lambda_{IR} = 2\pi c/\omega \) of the incident photons\textsuperscript{22-26}. This unique nano-IR apparatus has been routinely employed in studies of plasmon and phonon polaritons as well as for visualizing inhomogeneities in complex oxides\textsuperscript{19-21}. In the previous HPP
nano-imaging studies, a physical sample edge was chosen as the HPP wave reflector\(^5\), resulting in the imaging of coexisting fringes with two distinct periodicities \(\lambda_p\) and \(\lambda_p/2\). To avoid the complications of \(\lambda_p\) and \(\lambda_p/2\) fringe decompositions, an alternative approach is preferred. In our device (Figs.1b), pre-patterned micron-sized Au disk structures residing on top of the hBN crystals (bottom of \(\alpha\)-MoO\(_3\)), serve as fixed HPP antennas (Figs. 1,2) predominantly producing fringes with \(\lambda_p\) periodicity.

| Hyperbolic Bands (cm\(^{-1}\)) | hBN | \(\alpha\)-MoO\(_3\) |
|---------------------------------|-----|---------------------|
| \(\varepsilon_x = \varepsilon_y > 0, \varepsilon_z < 0\) | 760 - 820 | 545 - 850 |
| \(\varepsilon_x = \varepsilon_y < 0, \varepsilon_z > 0\) | 1365 - 1610 | 820 - 972 |

| Isotope | hB\(^{10}\)N | hB\(^{11}\)N | hBN | Nat. Ab. |
|---------|--------------|--------------|-----|---------|
| 98.7%   | 99.2%        | Nat. Ab.     |

Figure 1. Nano-IR imaging of phonon polaritons in hyperbolic strutures at cryogenic temperatures. a, Table with properties of examined hyperbolic materials b, Sketch of the layered Au/hBN and \(\alpha\)-MoO\(_3\)/Au structures. The lithographically-defined gold (Au) microstructures on top of the hBN (or below the \(\alpha\)-MoO\(_3\)) serve as effective plasmonic launchers. c, Nanoscale infrared image of phonon polariton fringes from the Au/h\(^{11}\)BN, expressed by the normalized scattering amplitude \(s\) acquired at a temperature of \(T = 50\) K. The arrows represent the propagation
direction of the plasmon waves. These experiments simultaneously visualize the local electric field associated with interference fringe patterns from phonon polaritons emitted by the Au microstructures.

We begin with a survey of a large-area (25 × 20 μm²) image of HPP standing waves in h¹¹BN obtained at T = 50 K (Fig. 1c) with an IR laser operating at λ IR = 6.6 μm. Here we display raw data in the form of the scattered near-field amplitude s normalized to the corresponding signal detected from the gold disks, whose optical response provides a convenient temperature (T)-independent reference. The most prominent aspect of the image in Fig. 1c is that the entire field of view is filled with HPP fringes. As expected, the λ p -fringes dominate the field of view, emanating from the Au antenna and propagating radially outwards. Even a cursory inspection of Fig. 1c reveals that HPP remain highly confined with λ IR/λ p > 20 and yet they travel over tens of microns, far exceeding previous measurements at ambient temperature⁶,¹³.

Nano-IR data in Fig. 2 attest to a clear reduction of HPP losses in monoisotopic hBN specimens at lower temperatures with the incident frequency of 1522 cm⁻¹ (6.57 μm). HPP in h¹¹BN films (thickness of 180 nm) at room temperature exhibit 15 μm propagation lengths, corresponding to quality factor Q p ~ 35 as defined below (Fig 2d&e). As the temperature is reduced, we observe a systematic increase in quality factor and propagation length. Specifically, at T = 45 K, Q p exceeds ~ 60 which represents the highest quality factor achieved to date. To elucidate the underlying polariton scattering mechanisms, we have also performed T-dependent studies in h¹⁰BN and naturally abundant samples for systematic comparisons (Fig. 2a&c). Two observations can be drawn. First, we observed that the monoisotopic specimens of h¹⁰BN and h¹¹BN share nearly identical propagation lengths at all temperatures. These HPP oscillations exhibit scattering lifetimes 2-3 times longer than those observed in naturally abundant hBN crystals, consistent with prior results⁶. Second, HPP propagation lengths at a given frequency
generally increase with reduced temperatures for all hBN specimens, as documented in Fig 2d&e. Notably, we found that both $h^{10}$BN and $h^{11}$BN exhibit a different temperature dependences compared with naturally abundant hBN crystals of similar thickness, as manifested by their HPP oscillations and the quality factor $Q_p$ (Fig 2d&e). These overall trends suggest different underlying mechanisms of phonon scattering between isotopically-pure and natural abundant hBN, as discussed in the following sections in detail.
Figure 2. Temperature dependence of phonon polariton propagation in isotopic hBN devices. 

a-c, Nano-IR images of the normalized scattering amplitude $s$ acquired at sequential sample temperatures from $h^{10}$BN, $h^{11}$BN and natural abundant hBN, respectively. A gold disk (labelled Au) functions as an antenna that emits hBN phonon waves. d, Line profiles of phonon polariton fringes propagating from left to right, as a function of the distance $L$ from the gold launcher. The variable attenuation of the propagating waveform (solid curves) is characteristic of the temperature-dependent phonon damping rate. Dash-dotted lines are fittings of the wave profile using damped sinusoidal functions corrected by geometric decaying factors (see main text and Methods). e, The temperature dependence of the quality factor $Q_p$ for phonon waves obtained from nanoscale infrared images shown in a-d. The thickness of $h^{10}$BN, $h^{11}$BN and naturally abundant hBN flakes are 150 nm, 200 nm and 180 nm, respectively.

For contrast, we have also investigated temperature dependence of hyperbolic HPP in exfoliated crystals of $\alpha$-MoO$_3$, an emerging natural material that exhibits in-plane hyperbolicity at mid-IR frequencies, to determine if the same temperature dependent scattering is observed as in hBN$^{13,14,16}$. In contrast to the convex wavefronts in isotropic materials, a concave wavefront of the HPP modes in thin slabs of $\alpha$-MoO$_3$ have been observed in ambient conditions$^{13,14,16,17}$. In Fig. 3, we show raster scans of an $\alpha$-MoO$_3$ crystals (≈ 200 nm thickness) exfoliated onto an Au antenna obtained at temperatures ranging from 50 to 300 K and at the frequency of 928 cm$^{-1}$. At room temperature, HPP interference fringes with concave shapes along the (100) direction were observed, consistent with the isofrequency curves as expected from the in-plane hyperbolic responses$^{13,14,16}$. As the temperature is reduced, we observe a systematic increase in both the overall propagation distance and the number of detectable HPP oscillations, consistent with hBN results. At $T = 50$ K, HPP oscillations approach propagation lengths of 20 µm (Fig. 3a). Similar temperature-dependence of HPP was also detected in the elliptical response regime (see SI). These overall trends are similar to our results on hBN isotopes, indicating the uniform characteristics of phonon propagation and dissipation emerging at low temperatures in hyperbolic crystals.
Figure 3. Temperature dependence of phonon polariton propagation in \( \alpha \)-MoO\(_3\) crystals. 

**a.** Nano-IR images of the normalized near-field scattering amplitude \( s \) acquired at sequential sample temperatures from \( \alpha \)-MoO\(_3\) crystals. A pair of gold disks beneath the \( \alpha \)-MoO\(_3\) crystal function as antennas that emit polaritonic waves. 

**b.** Line profiles of phonon polariton fringes propagating from left to right, as a function of the distance \( L \) from the gold launcher. Dash-dotted lines represent the results of numerical simulations performed to identify the temperature dependence of the complex
phonon polariton wavevector and associated damping rate (see main text and Methods). c, Temperature dependence of the quality factor $Q_p$ of phonon waves obtained from nanoscale infrared images shown in a.

To quantify the dynamics underlying propagation of HPP in hBN and $\alpha$-MoO$_3$ crystals, we present an analysis based on the complex momentum of the polariton $q_p = q'_p + i q''_p$. The real part defines the wavelength through $\lambda_p = 2\pi/q'_p$, whereas the imaginary part $q''_p$ quantifies dissipation and the quality factor $Q_p = q'_p/q''_p$. To quantify $Q_p$, we fit the oscillating line-profiles away from the Au antenna employing the spheroidal model with analytical formula of $S(x) = A \sin(q'_p x + B) e^{-q''_p x} / \sqrt{x+\zeta}$, with complex parameters $q_p$ and real $A$, $B$, and $\zeta$ as fitting parameters (see SI for details). This analytical formula accounts for fringe decay processes away from the antenna with the presence of a finite damping ($q''_p > 0$) and a geometric spreading term in the denominator. As shown by solid curves superimposed on polariton line-profiles in Fig. 2d & Fig. 3b, the fitted results match well with the experimental data across variable temperatures. This analysis allowed us to extract the temperature-dependent quality factors $Q_p(T)$ for polaritons in all isotopes of hBN and $\alpha$-MoO$_3$ crystals for the first time, as presented in Fig. 2e & Fig. 3c. Specifically, we found that the highest quality factor $Q_p(T)$ stems from the near-monoisotopic hBN flakes at cryogenic temperatures with $Q_p$ exceeding 60.

We now outline the damping rate analysis based on the temperature-dependent results extracted from hBN crystals. In Fig. 4, we plot the temperature-dependent HPP damping rates (linewidth) $\gamma(T) = \omega v_g/Q_p(T)v_p$ converted from quality factors, where $v_g$ and $v_p$ are the group and phase velocities of the modes. Note that the damping rates are on the order of several wavenumbers (approaching $\omega/1000$), while the quality factors defined in terms of momenta are
relatively small due to the very low group-phase velocity ratio: $\frac{v_g}{v_p} \sim 0.04$ (obtained from theoretical HPP dispersions, see SI). Several pieces of information arise from Fig. 4. Firstly, the HPP damping rates scale quasi-linearly with temperature for all samples. This is different from the nonlinear temperature dependence of plasmon damping rates observed in pristine graphene. Secondly, compared to h\textsuperscript{11}BN and h\textsuperscript{10}BN, the polariton damping rate in naturally abundant hBN displays a higher residual damping in the zero-Kelvin limit and a steeper slope to the temperature dependence.

The temperature dependent studies offer insights into the damping mechanisms of HPP in both hBN and \(\alpha\)-MoO\textsubscript{3} crystals. We posit that the temperature dependence can only be described by the participation of acoustic phonons in the scattering process, as the optical phonons are not thermally occupied in the temperature range of $T < 300 K = 208 \, cm^{-1}$ (see SI). In isotopically pure hBN, we attribute the leading damping channel to momentum relaxation caused by acoustic phonons, that is, the polariton being scattered from its initial state with momentum $k_i$ to the final state with momentum $k_f = k_i + q$ by absorbing/emitting an acoustic phonon with momentum $\pm q$ (Fig. 4b). Under reasonable approximations (see SI Sec. 4.1), the resulting scattering rate reads

$$\gamma_a(T) \sim \gamma_{a0}(\omega) f \left( \frac{T}{T_\Lambda} \right),$$

where $\gamma_{a0}(\omega)$ is a characteristic scattering rate, $T_\Lambda = v_a \Lambda/k_B$ is a temperature ‘cutoff’, $v_a$ is the speed of acoustic phonons and $\Lambda$ is the momentum cutoff of the hyperbolic polaritons. The temperature dependence is captured by the scaling function $f \left( \frac{T}{T_\Lambda} \right)$, which behaves as $1 + \frac{8n^4}{15} \left( \frac{T}{T_\Lambda} \right)^4$ when $T \ll T_\Lambda$ and as $1 + \frac{8}{3} \frac{T}{T_\Lambda}$ when $T \gg T_\Lambda$.

Isotopic impurities in natural hBN can also induce the same decay process as acoustic phonons, giving rise to a temperature-independent scattering rate $\gamma_i(\omega)$. Both $\gamma_i(\omega)$ and
\( \gamma_{a0}(\omega) \) are proportional to the polariton density of states \( D(\omega) = \partial_\omega \sqrt{-\frac{\varepsilon_x}{\varepsilon_z}} \), where \( \varepsilon_{x,z} \) are the in-plane/out-of-plane dielectric functions of hBN. Here \( D(\omega) \) diverges as \( \frac{1}{(\omega-\omega_{TO})^{3/2}} \) when the frequency approaches the transverse optical (TO) phonon, while weakly depending on \( \frac{1}{(\omega_{LO}-\omega)^{1/2}} \) as the frequency approaches the longitudinal optical (LO) phonons. Such a strong frequency dependence suggests that the frequency-dependent damping rate \( \gamma(\omega) \) needs to replace its constant counterpart in the Lorentzian oscillator model. Note that polaritons with frequency \( \omega \) can also decay into \( E_{1u} \) TO phonons or \( E_{2g} \) phonons (nearly flat dispersion at the frequency \( \omega_{TO} = \omega_g = 1360 \text{ cm}^{-1} \)) by emitting an acoustic phonon. This contributes to a scattering rate \( \gamma_g(\omega, T) = \gamma_{g0}(\omega)(n_b(\omega - \omega_{TO}) + 1) \) where the boson occupation number \( n_b \) gives the temperature dependence and \( \gamma_{g0} \) is a frequency scale depending on \( \omega \). However, our theoretical modeling suggests that this term is negligibly small (see SI, section 4). The functional forms of \( \gamma_a, \gamma_i \) and \( \gamma_g \) therefore represent these new results.

The steeper temperature-dependence of damping rate in natural hBN implies a higher-order decay channel involving both impurities and acoustic phonons, which we subsume by \( \gamma_h(T) = \kappa_i(\omega) \left( \gamma_a(\omega, T) + \gamma_g(\omega, T) \right) \) (see SI). The total polariton damping rates for natural and isotopically pure hBN are thus

\[
\gamma_{iso}(T) = \gamma_0 + \gamma_a(T) + \gamma_g(T), \quad \text{and} \quad \gamma_{nat}(T) = \gamma_{iso}(T) + \gamma_i + \gamma_h(T)
\]

where \( \gamma_0 \) is a temperature-independent scattering rate due to other decay processes\textsuperscript{27,28}. The dashed fitting lines in Figure 4 correspond to \( \gamma_0 = 0.4 \text{ cm}^{-1}, \gamma_{a0} = 0.6 \text{ cm}^{-1}, \gamma_{g0} = 0, \gamma_i = 0.5 \text{ cm}^{-1}, \kappa_i = 2.5 \) and \( T_\Lambda = 450 K \). Note that these fitting parameters are obtained for our simplified analytical model. Further dedicated first-principle calculations are possibly needed for
fine tuning, which is beyond the scope of the current work. On top of the above intrinsic damping channels, the dielectric loss from the SiO₂ substrate contributes < 20% to the total damping rate at the experimental frequency of 1522 cm⁻¹. Finally, the damping rate in α-MoO₃ is smaller than natural hBN, while slightly larger than monoisotopic hBN.

Figure 4. Theoretical modeling of phonon polariton damping rate $\gamma$ in hyperbolic crystals. (a) The dots corresponding to the extracted dissipation factors as a function of temperature; the solid blue & red lines are the theoretical fitting results as detailed in the main text. The parameters used here are: hB¹¹N: $\omega = 1522$ cm⁻¹, $v_g/v_p = 0.037$; hB¹⁰N: $\omega = 1553$ cm⁻¹, $v_g/v_p = 0.039$; naturally abundant hBN: $\omega = 1522$ cm⁻¹, $v_g/v_p = 0.039$; B¹¹: α-MoO₃ $= 928$ cm⁻¹, $v_g/v_p = 0.039$. The group and phase velocity in α-MoO₃ are taken from ref. [13] (b) The illustration of scattering of a hyperbolic polariton from an initial state (blue dot) to a final state (red dot) on the iso-frequency surface by either emitting/absorbing an acoustic phonon or by impurity potential.

In summary, our results account for HPP dynamics in high-quality hyperbolic crystals. Quality factors and scattering rates of hyperbolic polaritons in naturally abundant and monoisotopic hBN, as well as α-MoO₃ have been extracted from cryogenic nano-imaging studies. Though limited by acoustic phonon scattering and other decay channels, the HPP propagation length and lifetime in monoisotopic hBN at low-T exceed 25 μm and 5 ps, a new record among all natural hyperbolic materials reported so far. Our first report of HPP nano-imaging studies of high-
quality hyperbolic crystals at cryogenic temperatures demonstrates promising opportunities for the exploration of advanced phononic switching\textsuperscript{29}, polariton periodic orbits in cavities\textsuperscript{30} and nonlinear phenomena\textsuperscript{31} in ultra-pure polar samples at technologically relevant liquid nitrogen temperatures.

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Author contributions
All authors were involved in designing the research, performing the research, and writing the paper.

**Additional information**
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**Competing financial interests**
No competing financial interests.

**Data availability**
All other data that support the findings of this study are available from the corresponding author upon reasonable request.

**Code availability**
The custom code employed in this work to perform all calculations is available from the corresponding author upon reasonable request.

**Main Figure Legends**

**Methods**

1. **Sample growth and Device fabrication**

hBN crystals with the natural distribution of isotopes (20% B-10 and 80% B-11) were produced using a hot-pressed boron nitride ceramic boat, which served as both the container for the metal flux and as the B and N sources. For B$^{10}$ and B$^{11}$-enriched hBN crystal growth, the procedure is modified to use elemental boron as a source material. Therefore, the boron nitride boat was replaced with an alumina crucible. High-purity 10B (99.22 %) or 11B (99.41 %) powders were mixed with Ni and Cr powders to give overall concentrations of 4 % B, 48 % Ni and 48 % Cr. In this case, all nitrogen in the hBN originated from the flowing N$_2$ gas. Other than these changes, the procedure was the same as described above. The hBN crystals was then transferred onto SiO$_2$/Si substrates using the mechanical exfoliation method. After identifying the crystal thickness and uniformity, standard E-beam lithography is utilized to define Au launchers followed by thermal annealing processes to remove contaminated polymer residuals.

1. **Cryogenic IR Nano-Imaging Experiments**

IR nano-imaging experiments were performed using a home-built scattering-type scanning near-field optical microscope (s-SNOM) operating with variable sample temperatures from 50–300 K. All measurements are conducted under ultra-high vacuum conditions at pressures < 10$^{-9}$ mbar to prevent sample surface contamination. The cryogenic s-SNOM is equipped with continuous wave mid-IR quantum cascade lasers (daylightsolutions.com) and CO$_2$ lasers (accesslaser.com). The s-SNOM is based on an atomic force microscope (AFM), operated for the present experiments in non-contact mode using cantilevered metallic AFM probes with tip apex radius ~25 nm and tapping frequencies ~270 kHz. A pseudo-heterodyne interferometric detection
module is implemented in our s-SNOM to extract both scattering amplitude \( s \) and phase of the near-field signal. In the current work, we discuss only the former. In order to suppress background contributions to the back-scattered near-field signal, we demodulated the detected signal at the 3\(^{\text{th}}\) harmonic of the probe tapping frequency.