Chemical switching of low-loss phonon polaritons in α-MoO₃ by hydrogen intercalation

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Phonon polaritons (PhPs) have attracted significant interest in the nano-optics communities because of their nanoscale confinement and long lifetimes. Although PhP modification by changing the local dielectric environment has been reported, controlled manipulation of PhPs by direct modification of the polaritonic material itself has remained elusive. Here, chemical switching of PhPs in α-MoO₃ is achieved by engineering the α-MoO₃ crystal through hydrogen intercalation. The intercalation process is non-volatile and recoverable, allowing reversible switching of PhPs while maintaining the long lifetimes. Precise control of the intercalation parameters enables analysis of the intermediate states, in which the needle-like hydrogenated nanostructures functioning as in-plane antennas effectively reflect and launch PhPs and form well-aligned cavities. We further achieve spatially controlled switching of PhPs in selective regions, leading to in-plane heterostructures with various geometries. The intercalation strategy introduced here opens a relatively non-destructive avenue connecting infrared nanophotonics, reconfigurable flat metasurfaces and van der Waals crystals.
Polaronics are hybrid electromagnetic waves originating from the coupling of light and dipole-like excitations in matter and include phonon polaritons (PhPs), plasmon polaritons, and exciton polaritons. Among these polaritonic modes, PhPs with strong field confinement, low loss, and a long lifetime can enable a wide range of applications in light transmission, electro-optical modulation, sub-diffraction focusing, and imaging, and molecular detection. However, to best utilize PhPs, external control of their properties is needed, which is a difficult task owing to the wide bandgap (~2–10 eV) in polar crystals, making the common electrical-gating method impractical.

Although a few attempts have been made, effective tuning of PhPs remains challenging. Recently, PhP manipulation has been investigated by controlling the dielectric environment through the fabrication of vertical heterostructures of van der Waals (vdW) crystals (hexagonal boron nitride (h-BN)/black phosphorous11 and MoS₂/SiC12), polaritonic metasurfaces in combination with phase change materials (h-BN/VO₂13), germanium antimony telluride (GeSbTe)/quartz16 and h-BN/GeSbTe17), or polar materials suspended in air18. The latter combination with phase change materials (h-BN/VO₂) which different characterizations of the as-prepared exfoliation process (see Methods and Supplementary Fig. 1, in vapor deposition (CVD) method, followed by a mechanical exfoliation process (see Methods and Supplementary Fig. 1, in which different characterizations of the as-prepared α-MoO₃ flakes confirm their orthorhombic crystal structure). The hydrogen intercalation process was conducted by treating the flakes with a hydrogen plasma. To avoid irreversible damage to the crystal, the exposure time was controlled at 10 s (see Methods and Supplementary Fig. 2). Four typical hydrogen molybdenum bronzes (HₓMoO₃, 0 < x ≤ 2) with different compositions and structures can be obtained after α-MoO₃ hydrogenation. In our experiment, we obtained type I HₓMoO₃ (orthorhombic, 0.2 < x < 0.4) in hydrogenated α-MoO₃ (hereafter, termed H-MoO₃), as revealed by Raman and X-ray diffraction (XRD) analysis. In the MoO₃ octahedron, the oxygen atoms can be divided into three types according to their site (Fig. 1a). As shown in Fig. 1b, c, the Raman shifts at 667, 818, and 994 cm⁻¹ are assigned to the stretching vibration modes of the Mo–O₃, Mo–O₂, and Mo–O₁ bonds, respectively. After hydrogenation, owing to a structural rearrangement, the peaks at 818 and 994 cm⁻¹ shift to 760 (broadband) and 1008 cm⁻¹ (black arrows in Fig. 1b), respectively, agreeing well with the Raman spectrum of type I HₓMoO₃.28 Another stretching vibration peak of the Mo–O₁ bond at 441 cm⁻¹, which is hardly observable in O-MoO₃, emerges after hydrogenation. Given that the Raman peaks at 667, 818, and 994 cm⁻¹ remain discernible in H-MoO₃, even those with weaker intensities (Supplementary Fig. 3), the hydrogenated product is a mixture of type I HₓMoO₃ and α-MoO₃. In addition, the XRD spectra of O-MoO₃ (Fig. 1d,e) correspond to the Pmmn space group of orthorhombic α-MoO₃ (ICDD PDF: 05-0508) with lattice constants a = 3.97 Å, b = 13.86 Å, and c = 3.71 Å. After hydrogenation, new peaks arise at 12.6°, 25.3°, and 38.3° (black arrows), which are attributed to the (020), (040), and (060) planes of orthorhombic HₓMoO₃ (ICDD PDF: 70-0615). This result unambiguously proves the formation of type I HₓMoO₃ with calculated lattice constants a = 3.93 Å, b = 14.02 Å, and c = 3.74 Å.

The as-prepared H₂MoO₃ is stable at room temperature and can last for several months in the ambient atmosphere (Supplementary Fig. 4). However, we can apply a simple thermal treatment (see Methods) to induce deintercalation, which proceeds as

\[ \text{H}_\text{x}\text{MoO}_3 + \text{O}_2 \rightarrow \text{MoO}_3 + \text{H}_2\text{O} \] (1)

Both the Raman and XRD spectra of the recovered α-MoO₃ (hereafter, termed R-MoO₃) agree excellently with those of O-MoO₃, demonstrating that the H-MoO₃ crystal structure is reconfigured to α-MoO₃ after dehydrogenation. Moreover, X-ray photoelectron spectroscopy (XPS) characterizations (Supplementary Fig. 5 and Supplementary Table 1) provide solid evidence for the reversibility of the chemical composition changes that occur during the intercalation/deintercalation process. Compared with the metal-atom intercalation, our approach thus affords an effective and relatively non-destructive strategy to reversibly modify the compositional structure and property of nanometer-thick transition metal oxide flakes.

Reversible switching of PhPs. The structural reversibility during hydrogenation/dehydrogenation processes enables direct manipulation of the vibration-dependent PhPs in nanometer-thick α-MoO₃ slabs. To study the polaritonic response of our samples, we used scattering-type scanning near-field optical microscopy (s-SNOM, Fig. 2a), which enables direct mapping of the near field in terms of the amplitude and phase and the sample topography by illuminating the sample with mid-infrared light at different frequencies. First, we measured the samples in the lower Reststrahlen band (1-RB, 818–974 cm⁻¹) of α-MoO₃, where PhPs are known to propagate with a hyperbolic wavefront along the [100] direction. Figure 2b shows s-SNOM amplitude images (third-harmonic signals, s₃) at frequency ω = 890 cm⁻¹ together

Results

Structural analysis during reversible hydrogenation. As a typical vdW material, α-MoO₃ is composed of double layers of linked and distorted MoO₃ octahedra, as shown in Fig. 1a. The nanometer-thick α-MoO₃ (original α-MoO₃, hereafter termed O-MoO₃) flakes studied in this work were synthesized by a chemical vapor deposition (CVD) method, followed by a mechanical exfoliation process (see Methods and Supplementary Fig. 1, in which different characterizations of the as-prepared α-MoO₃...
**Fig. 1 Structural characterizations during the hydrogenation/dehydrogenation process.**

- **a** Schematic illustrations of the crystal structures (linked double layers) and bond lengths (Å, ref. 28) of \( \alpha \)-MoO\(_3\) and type I H\(_x\)MoO\(_3\). O\(_1\), O\(_2\) and O\(_3\) represent terminal oxygens, doubly coordinated bridging oxygens and triply coordinated oxygens, respectively.
- **b** Raman spectra of the original (O-MoO\(_3\)), hydrogenated (H-MoO\(_3\)), and recovered (R-MoO\(_3\)) \( \alpha \)-MoO\(_3\). Black arrows indicate the vibration peaks that emerged after hydrogenation.
- **c** Magnified Raman spectra (normalized) of O-MoO\(_3\), H-MoO\(_3\), and R-MoO\(_3\) in **b**.
- **d** XRD pattern of the original, hydrogenated and recovered \( \alpha \)-MoO\(_3\). Red and green tick marks represent the peaks in ICDD PDF card no. 05-0508 and 70-0615, respectively. Black arrows indicate the diffraction peaks originating from type I H\(_x\)MoO\(_3\).
- **e** Comparison of the (020), (040), and (060) diffraction peaks (normalized) in **d**.

**Fig. 2 Reversible switching of PhPs in the L-RB of \( \alpha \)-MoO\(_3\).**

- **a** Schematic of the s-SNOM measurement and PhP propagation in a typical H-MoO\(_3\)/\( \alpha \)-MoO\(_3\) in-plane heterostructure.
- **b** Optical micrographs and s-SNOM amplitude images (890 cm\(^{-1}\)) of a MoO\(_3\) flake on a SiO\(_2\)/Si substrate before intercalation, after intercalation (10 s) and after deintercalation. Scale bar = 5 \( \mu \)m.
- **c** PhP dispersion data and fitting results (solid lines) along the [100] direction. The error bars define the 95% confidence intervals.
- **d** Group velocities of PhPs in original and recovered flakes extracted from the fitting results in **c**.
- **e** Extracted real-part near-field line traces and fitting curves (solid lines) at 903 cm\(^{-1}\).
- **f** Comparison of the group velocities and lifetimes of PhPs (903 cm\(^{-1}\)) after three reversible intercalation cycles. The error bars of lifetimes were calculated based on the errors of propagation lengths originating from the standard deviation of fitting.
with optical micrographs of O-MoO$_3$, H-MoO$_3$, and R-MoO$_3$ flakes with the thicknesses of 220 nm. As expected, for O-MoO$_3$, we observe interference fringes in the s-SNOM image, revealing the excitation of PhPs propagating parallel to the [001] direction with a polariton wavelength $\lambda = 1.61 \pm 0.01$ µm, which well matches previous work$^{19}$. After intercalation with hydrogen (H-MoO$_3$), we observe a weaker contrast in the optical image, suggesting a varied optical behavior. More importantly, no regular interference fringes can be observed in the s-SNOM image (Supplementary Fig. 6), implying that PhPs are not excited. After dehydrogenation, the initial contrast of the flake in the optical image is recovered, and parallel fringes in the s-SNOM image are again observed, revealing the excitation of PhPs.

For a quantitative analysis of the PhP behavior, we compared the group velocities ($v_g$), propagation lengths ($L$), and lifetimes ($\tau$) of PhPs in the original and recovered α-MoO$_3$ flakes. First, we extracted the PhP dispersions ($\omega(Re(k))$, where $k$ is the complex-valued wavevector) along the [100] direction (dashed lines in Fig. 2b) at different frequencies by fitting the line scans obtained from s-SNOM amplitude and phase images using the function$^{29}$:

$$\sigma(x) = A \frac{e^{2kx}}{\sqrt{x}} + B \frac{e^{kx}}{x^d}$$

(2)

where $\sigma$ is the complex-valued optical signal, $d (~1)$ is variable decay, $A$ and $B$ are the parameters for tip- and edge-launched PhPs, respectively. We used the function $y = ax^d$ to fit the dispersion data (solid lines in Fig. 2c, see Supplementary Table 2) and then performed a numerical derivation process on the fitting curve, as the group velocity can be extracted by $v_g = \partial \omega / \partial Re(k)$.$^{19}$ The resulting group velocities of PhPs in the original and recovered α-MoO$_3$ flakes are shown in Fig. 2d. We observe similar values of $4.42 \times 10^{-3}$ c and $4.45 \times 10^{-3}$ c at $\omega = 903$ cm$^{-1}$ for the original and recovered flakes, respectively, where $c$ is the speed of light in vacuum. Then we calculate the polariton lifetime by $\tau = \nu / v_g$ (see Supplementary Note 1 for more details). Accordingly, the line profiles at $\omega = 903$ cm$^{-1}$ and the fitting results are plotted in Fig. 2e, where the propagation lengths of PhPs in the original and recovered α-MoO$_3$ are $2.14 \pm 0.61$ µm and $2.09 \pm 0.22$ µm, respectively. The calculated lifetime of PhPs in O-MoO$_3$ (1.61 ± 0.46 ps) is comparable to that in R-MoO$_3$ (1.57 ± 0.16 ps), suggesting high-fidelity recovery of hyperbolic PhPs after dehydrogenation. The PhP behavior remains almost unchanged even after three intercalation/deintercalation cycles, as shown in Fig. 2f (Supplementary Fig. 7).

In addition to PhPs in the L-RB, PhPs with elliptical propagation in the upper Reststrahlen band (U-RB, 960–1010 cm$^{-1}$) of α-MoO$_3$ can also be reversibly switched through hydrogenation while maintaining their extraordinary properties (strong field confinement and ps lifetimes$^{19}$). As shown in Fig. 3a, PhPs propagate along both the [100] and [001] directions in O-MoO$_3$ and R-MoO$_3$ owing to the elliptical PhP propagation. We compared the performances of PhPs in the original and recovered α-MoO$_3$ flakes (140 nm) along the two directions (Fig. 3b-d). In the [100] direction (dashed line 1 in Fig. 3a), the PhP group velocities and lifetimes are $0.36 \times 10^{-3}$ c and $10.93 \pm 2.96$ ps (original) and $0.35 \times 10^{-3}$ c and $10.38 \pm 1.71$ ps (recovered) at 990 cm$^{-1}$, respectively. Comparable characteristics are also found in the [001] direction (dashed line 2), where the group velocities (lifetimes) in the original and recovered slabs are $0.43 \times 10^{-3}$ c ($12.25 \pm 1.94$ ps) and $0.42 \times 10^{-3}$ c ($12.86 \pm 1.67$ ps), respectively. These results demonstrate that the hydrogen intercalation/deintercalation method enables robust, permanent (several months) and reversible switching of PhPs in α-MoO$_3$ both in the L- and U-RB.

**Analysis of intermediate states during hydrogenation.** To study the hydrogen intercalation process and the evolution of the PhPs excited in nanometer-thick α-MoO$_3$, we prepared H-MoO$_3$ flakes with various hydrogen proportions by precisely controlling the intercalation time (i.e., hydrogen plasma treatment time, Supplementary Fig. 8). The XPS results (Fig. 4a and Supplementary Table 3) show a slowly increasing proportion of Mo$^5+$ (H$_x$MoO$_3$) with prolonged intercalation time. After intercalation, we used s-SNOM to investigate the effect on the PhP properties in these samples. Figure 4b shows normalized s-SNOM amplitude (top) and phase (bottom) images of a 120 nm-thick α-MoO$_3$ flake before (O-MoO$_3$ (0 s)) and after 2, 5, 7, and 10 s hydrogen intercalations (H-MoO$_3$). The parallel interference fringes in both

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**Fig. 3** Reversible switching of PhPs in the U-RB of α-MoO$_3$. **a** Optical micrographs and s-SNOM amplitude images (990 cm$^{-1}$) of original, intercalated, and recovered MoO$_3$ flakes. Scale bar = 5 µm. **b** PhP dispersion data and fitting results (solid lines) along both [100] and [001] directions (dashed lines in **a**). The error bars represent the 95% confidence intervals. **c** Group velocities of PhPs extracted from the fitting results in **b**. **d** Comparison and fitting (solid lines) of extracted line traces.
the amplitude and phase images reveal that the excitation of PhPs gradually vanishes with increasing intercalation time, which can be further verified by the extracted amplitude line traces (Fig. 4c) and fast Fourier transform (FFT) results (Supplementary Fig. 9). The quantitative comparison indicates that upon gradual hydrogenation, the PhP dispersion has no obvious difference, while the propagation length and lifetime gradually decrease. By comparing the Raman spectra in hydrogenated intermediate states, we attribute this chemical switching of PhPs to the disappearance of optical phonons (L-RB and U-RB) of the polar material during hydrogenation (Supplementary Note 2 and Supplementary Fig. 10).

Interestingly, we found regular acicular nanostructures in intermediate H-MoO$_3$ (2 and 5 s) compounds. For a better analysis, we prepared an intercalated flake (2 + 2 s) by a step-by-step method (Supplementary Fig. 11). We observe needle-like nanostructures (Fig. 4d) with a weaker optical contrast extending along ~57 ± 2° with respect to the [100] direction (on the (010) surface), which can be assigned to the <203> direction. Similar nanostructures are also observed in nanometer-thick α-MoO$_3$ discs (Supplementary Fig. 12), indicating that the hydrogenation direction determined by the intrinsic crystal structure is the main reason for their nucleation orientation instead of the specific geometry of the flake. Indeed, by performing Raman intensity mapping, we discover that these nanostructures consist of type I H$_x$MoO$_3$ (Supplementary Fig. 12).

To better understand the nucleation of these nanostructures, we studied the diffusion of hydrogen atoms in type I H$_x$MoO$_3$ by performing calculations using density functional theory (DFT) (see Methods and Supplementary Note 3). The O$_2$ sites (with an adsorption energy of 3.58 eV) are the most energetically favorable
for hydrogen adsorption compared with the O1 (3.06 eV) and O3 (1.13 eV) sites (Supplementary Table 4). In addition, hydrogens are alternately located along the zig-zag chains of intralayer oxygen atoms ([001] direction) within MoO3 octahedra. By comparing the calculated thermodynamic stabilities for hydrogen diffusion along the [003] direction and along the [001] direction, we find the former to be slightly more energetically favorable (15 meV), possibly because of the cationic repulsion (Supplementary Fig. 13). We can thus conclude that the three-dimensional hydrogen diffusion in H-MoO3 at low hydrogen concentration occurs along the [003] direction, as schematically shown in Fig. 4e, giving rise to the observed needle-like nanostructures oriented along 57 ± 2° with respect to the [100] direction.

Further characterization by s-SNOM of these nanostructures shows that while in terms of the topography (Fig. 4d and Supplementary Fig. 14), they are relatively hard to identify (probably because of their atomic expansion of ~4 nm caused by the rearrangement of MoO6 octahedra, which can be recovered after deintercalation), in the amplitude image, they show a strong near-field contrast compatible with a high degree of metallicity because of the greatly enhanced conductivity of H-MoO3 (Supplementary Fig. 15). Clear parallel interference fringes revealing excitation and/or reflection of PhPs are observed on both sides of the nanostructure (Fig. 4f). In fact, by extracting amplitude line scans perpendicular to the [003] direction (i.e., [302]) between two parallel nanostructures (Fig. 4g) in the red rectangular region in Fig. 4d, we obtain resonator-like oscillations with frequency-locked wavelengths similar to PhPs along that direction. We also compare λp at different frequencies in O-MoO3, R-MoO3, and four typical regions in H-MoO3 (rectangles in Fig. 4d). As displayed in Fig. 4h, the λp of PhPs along the [100] direction remains similar in O-MoO3, R-MoO3, and H-MoO3. However, PhPs along the <203> direction in H-MoO3 exhibit decreased polariton wavelengths, which are much shorter than those of PhPs along the same direction in O-MoO3. The FFT results along the [302] direction in Supplementary Fig. 16 show two prominent peaks, which is attributed to PhPs launched by both the tip and acicular nanostructures; thus, we can conclude that these hydrogenated nanostructures act as efficient reflectors and launchers of PhPs (see Supplementary Fig. 17 for U-RB) while forming part of the crystal (as defined chemically and not physically), which can be highly beneficial for the development of planar devices in nanophotonic technologies (Supplementary Note 4).

Spatially controllable switching of PhPs. Finally, we studied the scalability and feasibility of spatially selective hydrogenation within α-MoO3 flakes. As illustrated in Fig. 5a, we combined the hydrogenation method with a conventional photolithography technique. We used direct photolithography to cover a 225 nm-thick α-MoO3 flakes by a photore sist with designed patterns (see Methods and Supplementary Fig. 18 and 19 for details). In this way, upon hydrogenation, only open regions of α-MoO3 flakes are exposed, and thus, H-MoO3/α-MoO3 in-plane heterostructures with various geometries can be achieved. Figure 5b, c shows a flake where the bottom half region has been hydrogenated and thus PhPs are switched off in both the L-RB and U-RB. In contrast, in the hydrogen-free region (α-MoO3), parallel interference fringes indicating PhPs are observed.

Owing to its in-plane anisotropy, α-MoO3-based in-plane nanostructures with different orientations are attractive for nanoptics studies and can be easily prepared using our method by designing the masked region as shown in Fig. 5d, where the boundary of the heterostructure is perpendicular to the PhP propagation direction. Furthermore, the thickness of the hydrogenated regions shows no obvious difference from that of the unhydrogenated regions, whereas the optical signal is very different, as confirmed by the extracted line traces (Fig. 5d). Figure 5e shows the potential application of our chemical switching method. By using customized patterns in photolithography, discs, squares, and more complicated arrays with various PhP dispersions can be constructed. Notably, we observe a clear reflection of PhPs at the sharp boundaries along the <203> direction (Fig. 4d and square in Fig. 5e). However, no significant reflection occurs at the boundaries
along other directions (Fig. 5d and disc in Fig. 5e) because these boundaries are composed of much smaller \(<20\text{3}^-\) oriented mer40, programmable in-plane heterostructures and magnetic properties35, and high conductivity of hydrogen-doped \(\alpha\)-MoO\(_3\) flakes. We also achieve spatially controllable intercalation and selective switching of PhPs by combining our intercalation process with photolithography (or possibly a shadow mask), through which in-plane heterostructures are constructed, which is difficult to realize in other PhP tuning methods (Supplementary Note 5 and Supplementary Table 5).

Our methodology establishes a proof of concept for chemically manipulating polaritons, offering opportunities for the growing nanophotonics community. We envision that in operando switching of PhPs could be realized during s-SNOM nanoimaging via tip-induced hydrogen intercalation (the so-called spillover technique33). The plasmonic resonance34, newly observed ferromagnetic properties35, and high conductivity of hydrogen-doped MoO\(_3\) enable analysis of the interaction of PhPs with plasmon and magnetic fields. Furthermore, using other intercalators, such as ions (Li\(^+\), Na\(^+\))36,37, metal atoms (Co, Fe)38,39, and polymers40, programmable in-plane heterostructures and flat meta-surfaces could be constructed, and applications combining scalable planar optics with photomagnetics, photoelectronics and topological photonics could be realized.

**Methods**

**Preparation of \(\alpha\)-MoO\(_3\) flakes.** \(\alpha\)-MoO\(_3\) bulk crystals were grown by a CVD method. In brief, 0.1 g of MoO\(_3\) particle (Alfa Aesar) was placed in the middle of a quartz tube (2 inches in diameter), where the temperature was 780 °C. Ar (200 standard cubic centimeters per minute (scm)) and O\(_2\) (50 scm) were used as carrier gases to reduce the oxygen vacancies. \(\alpha\)-MoO\(_3\) bulk crystals were collected at the down-stream position where the temperature was ~590 °C. Nanometer-thick \(\alpha\)-MoO\(_3\) flakes were mechanically exfoliated and then transferred to a SiO\(_2\) (300 nm)/Si substrate.

**Reversible hydrogen intercalation/deintercalation processes.** The hydrogen intercalation process was conducted by a plasma-enhanced CVD method. Typically, \(\alpha\)-MoO\(_3\) flakes on a SiO\(_2\)/Si substrate (1 \times 1 cm\(^2\)) were placed in the middle of a quartz tube (four inches in diameter), where the temperature was 150 °C. A radio-frequency plasma generator (17 mW, 13.56 MHz, VERG-500) was placed upstream of the sample at a distance of ~50 cm. H\(_2\) (5 scm) was used as the gas source under a pressure of 15 mTorr. By controlling the reaction time, hydrogen-intercalated flakes with various hydrogen levels were obtained. The deintercalation experiment was carried out by a thermal treatment process at 350 °C for 120 min. Ar (200 scm) and O\(_2\) (50 scm) were used as carrier gases at atmospheric pressure to reconfigure the crystal structure and reduce oxygen defects. Notably, to obtain H\(_x\)MoO\(_3\) flakes with different hydrogen distributions, two intercalation strategies were carried out: continuous intercalation and step-by-step intercalation.

**Spatially controllable hydrogen intercalation process.** Spatially controlled hydrogenation in nanometer-thick \(\alpha\)-MoO\(_3\) flakes was realized by combining intercalation with photolithography. First, a layer of photore sist (~1 μm, AZ 1512, AZ Electronic Materials) was coated on top of the surfaces of \(\alpha\)-MoO\(_3\) flakes and then patterned by direct-write lithography (SF100 XPRESS). After the development process, the unexposed regions of \(\alpha\)-MoO\(_3\) flakes were covered and protected by the photore sist. Then, a similar hydrogenation process was conducted, during which hydrogen was inserted into the exposed regions of flakes. Following a lift-off process, selectively intercalated flakes (i.e., in-plane H:MoO\(_3\)/\(\alpha\)-MoO\(_3\) heterostructures) with designed geometries were obtained.

**Theoretical calculations.** We investigated the adsorption of hydrogen on \(\alpha\)-MoO\(_3\) using DFT calculations as implemented in the Vienna ab initio simulation package. The Perdew–Burke–Ernzerhof form of the generalized gradient approximation was used to describe electron exchange and correlation. The plane-wave kinetic energy cutoff was set to 600 eV. A semiempirical function developed by Grimme (DFT-D3) was employed to describe dispersion forces. All structures were relaxed until the ionic forces were <0.01 eV Å\(^{-1}\). For accurate calculations of the electronic structures, we used 9×3×9 \(\Gamma\) centered grids for sampling the Brillouin zone to calculate the preferred adsorption site of one hydrogen on \(\alpha\)-MoO\(_3\), and 5×5×3 \(\Gamma\) centered grids to simulate the hydrogen diffusion process in a 6×1×6 \(\alpha\)-MoO\(_3\) supercell.

**Characterizations.** Optical observations and Raman tests were conducted using a confocal microscope system (WITec, alpha 300 R). TEM and SAED images were obtained from a FEI Tecnai T20. HAADF-STEM images were acquired by a JEOL JEM-ARM300F TEM. XRD and XPS spectra were collected by a Bruker D8 Advance and a Nexsa surface analysis system (Thermo Scientific), respectively. Surface topography and near-field images were captured using a commercial s-SNOM (Nearfield Imaging, Arrow-NCPI) in tapping mode (~270 kHz frequency and ~70 nm amplitude). To eliminate edge-excited polaritons, flakes were rotated till the incident polarization was parallel to the edges.

**Data availability**

The data that support the findings of this study are available from the corresponding authors upon reasonable request.

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**References**

1. Basov, D. N., Fogler, M. & de Abajo, F. G. Polaritons in van der Waals materials. Science 354, aaa9922 (2016).
2. Fei, Z. et al. Gate-tuning of graphene plasmons revealed by infrared nanoimaging. Nature 487, 82–85 (2012).
3. Dai, S. et al. Tunable phonon polaritons in atomically thin van der Waals crystals of boron nitride. Science 343, 1125–1129 (2014).
4. Hu, F. et al. Imaging exciton–polariton transport in MoSe\(_2\) waveguides. Nat. Photon. 11, 356–360 (2017).
5. Low, T. et al. Polaritons in layered two-dimensional materials. Nat. Mater. 16, 182–194 (2017).
6. Sun, Z., Martinez, A. & Wang, F. Optical modulators with 2D layered materials. Nat. Photon. 10, 227–238 (2016).
7. Li, P. et al. Hyperbolic phonon polaritons in boron nitride for near-field optical imaging and focusing. Nat. Commun. 6, 7305 (2015).
8. Autore, M. et al. Boron nitride nanoresonators for phonon-enhanced molecular vibrational spectroscopy at the strong coupling limit. Light Sci. Appl. 7, 17172 (2018).
9. Dunkelberger, A. D. et al. Active tuning of surface phonon polariton resonances via carrier photoinjection. Nat. Photon. 12, 50–56 (2017).
10. Giles, A. J. et al. Ultralow-loss polaritons in isotopically pure boron nitride. Nat. Mater. 17, 134–139 (2018).
11. Chaudhary, K. et al. Engineering phonon polaritons in van der Waals heterostructures to enhance in-plane optical anisotropy. Sci. Adv. 5, eaau7117 (2019).
12. Dubrovkin, A. M., Qiang, B., Krishnamoorthy, H. N. S., Zheludev, N. I. & Vlcek, A. A. M. Biaxial hyperbolic van der Waals crystal. Nat. Commun. 10, 4887 (2019).
13. Folland, T. G. et al. Reconfigurable infrared hyperbolic metasurfaces using phase change materials. Nat. Commun. 9, 4371 (2018).
14. Dai, S. et al. Phase-change hyperbolic heterostructures for nanopolaritons: A case study of hBN/VO\(_2\). Adv. Mater. 31, 1900251 (2019).
15. Fals, A. et al. Receptive index-based control of hyperbolic phonon–polariton propagation. Nano Lett. 19, 7725–7734 (2019).
16. Li, P. et al. Reversible optical switching of highly confined phonon–polaritons with an ultrathin phase-change material. Nat. Mater. 15, 870–875 (2016).
17. Chaudhary, K. et al. Polariton nanophotonics using phase change materials. Nat. Commun. 10, 4887 (2019).
18. Dai, S. et al. Hyperbolic phonon polaritons in suspended hexagonal boron nitride. Nano Lett. 19, 1009–1014 (2019).
19. Ma, W. et al. In-plane anisotropic and ultra-low-loss polaritons in a natural van der Waals crystal. Nature 562, 557–562 (2018).
20. Zheng, Z. et al. A mid-infrared biaxial hyperbolic van der Waals crystal. Sci. Adv. 5, eaav8690 (2019).
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Author contributions

Y.W. and Q.O. contributed equally to this work. Q.B. conceived the original concept. G.L., X.L., and Q.B. supervised the project. Y.W. and Q.O. designed and conducted most of the experiments, analyzed the data and prepared the manuscript. Y.Y. under the supervision of N.M. calculated the hydrogen diffusion pathway. Y.L. and W.Y. conducted the photolithography experiment. W.M., Z.D., and B.S. contributed to the s-SNOM measurements. X.C. and X.B. contributed to sample preparation. J.D., G.A.-P., and P.A.-G. contributed to the analysis of phonon polaritons. Y.W., Q.O., G.L., X.L., C.-M.L., P.A.-G., and Q.B. participated in data analysis and co-wrote the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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