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Multi-resonance enhanced photothermal synergistic fiber-optic Tamm plasmon polariton tip for high-sensitivity and rapid hydrogen detection

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Accurate and real-time detection of hydrogen (H₂) is essential for ensuring energy security. Fiber-optic H₂ sensors are gaining attention for their integration and remote sensing capabilities. However, they face challenges, including complex fabrication processes and limited response times. Here, we propose a fiber-optic H₂ sensing tip based on Tamm plasmon polariton (TPP) resonance, consisting of a multilayer metal/dielectric Bragg reflector deposited directly on the fiber end facet, simplifying the fabrication process. The fiber-optic TPP (FOTPP) tip exhibits both TPP and multiple Fabry-Perot (FP) resonances simultaneously, with the TPP employed for highly sensitive H₂ detection. Compared to FP resonance, TPP exhibits more than twice the sensitivity under the same structural dimension without cavity geometry deformation. The excellent performance is attributed to alterations in phase-matching conditions, driven by changes in penetration depth of TPP. Furthermore, the FP mode is utilized to achieve an efficient photothermal effect to catalyze the reaction between H₂ and the FOTPP structure. Consequently, the response and recovery speeds of the FOTPP tip under resonance-enhanced photothermal assistance are improved by 6.5 and 2.1 times, respectively. Our work offers a novel strategy for developing TPP-integrated fiber-optic tips, refines the theoretical framework of photothermal-assisted detection systems, and provides clear experimental evidence.

Keywords: fiber-optic hydrogen sensor; Tamm plasmon polariton; photothermal synergistic effect; dynamic response enhancement; cost-effective production

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Introduction

Hydrogen (H_2) , as a green renewable energy source, is anticipated to replace traditional fossil fuels in industries,

construction, transportation, and other fields due to its excellent combustion performance and environmentally friendly characteristics^{1,2}. However, H_2 is a highly

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flammable and explosive gas, with a risk of explosion when its concentration in the air exceeds 4%. Additionally, H₂ is colorless and odorless, difficult to perceive directly. Therefore, it is extremely crucial to develop highperformance H₂ sensors for real-time detection of H₂ concentration in these processes involving its production, storage, transportation, and utilization³. Optical sensors based on hydrogen-sensitive materials like palladium (Pd), platinum (Pt), and their alloys, have become prominent and feasible solutions for real-time detection^{4–8}. Unlike traditional electrical sensors^{9,10}, these optical sensors exhibit a reduced risk of explosion and lower power consumption, making them more competitive for various application scenarios. Moreover, with the rapid development of Internet of Things (IoT) technology, optical sensors are increasingly focused on miniaturization and integration.

Fiber optic sensors, as integrated and miniaturized alternatives to prism or on-chip devices, combine optical sensing with optical transmission capabilities. They offer enhanced flexibility, robust resistance to electromagnetic interference, and the potential for remote monitoring and distributed measurement¹¹⁻¹⁸. Consequently, considerable efforts have been devoted to developing highly sensitive and fast-response fiber-optic H₂ sensors using various resonance coupling mechanisms, such as Fabry-Perot (FP) interferometers^{19,20}, fiber Bragg gratings^{21,22}, tilted fiber Bragg gratings²³, Mach-Zehnder interferometers²⁴, fiber optic surface plasmon resonance^{25,26}, and the like. The response time of these fiber-optic sensors typically depends on the H₂ permeability of the H₂-sensitive material, which is negatively correlated with its thickness and positively correlated with the working temperature²⁷. Therefore, increasing the operating temperature is an effective strategy to improve the response time of the H₂ sensor with fixed structural dimensions. Contemporarily, photothermal effects have been integrated into various emerging optoelectronic devices for their precise and non-contact heating capabilities²⁸⁻³³. This approach provides an all-optical pathway to develop temperatureassisted fiber-optic sensors for rapid H₂ detection^{34,35}. Nevertheless, the theoretical analysis and experimental investigation of the photothermal effect in the reported photothermal-assisted H₂ detection remain insufficient, hindering the optimal utilization of the photothermal effect in these systems.

Despite significant advancements in developing highperformance fiber-optic H₂ sensors have been achieved,

these efforts often rely on complex fabrication processes such as photolithography, fiber modification, and structural transfer, which restricts the potential for achieving low-cost and large-scale production of fiber-optic H₂ sensors. Tamm plasmon polariton (TPP), as a boundary state mode, is excited at the interface between the distributed Bragg reflector (DBR) and metal film³⁶⁻³⁹. The geometry structure supporting the TPP mode is constructed using only planar multilayer films, bypassing the requirement for photolithography or structural transfer methods. Consequently, TPP in multilayer films has attracted considerable attention in various fields, including biochemical sensing^{40,41}, filtering and absorption^{42,43}, nonlinear optics44, and electro-optic modulation45. However, electric fields in TPP-based optical sensors are mainly concentrated inside the structure, resulting in relatively low sensitivity and thus not appropriate for bioassays. One effective solution is to replace the metal film with sensitive materials to achieve high-sensitivity detection of specific analytes⁴⁰, which is consistent with the principle of H₂ detection⁴⁶. Furthermore, the excitation of TPP does not necessitate strict control of the angle and polarization of incident light. Therefore, it can be directly integrated into the fiber end facet without requiring special fiber treatment. Nevertheless, to the best of our knowledge, TPP-integrated fiber-optic devices constructed by multilayer planar films, including sensors and other optoelectronic devices, have not been experimentally reported until now.

In this paper, we experimentally integrate multilayer planar DBR and H2-sensitive metal Pd onto the end facet of fiber for the first time, supporting the excitation of TPP and FP modes. By assigning distinct functions to different resonance modes, a multi-resonance enhanced photothermal synergistic FOTPP H₂ sensing tip is both theoretically and experimentally demonstrated. In this system, TPP mode with higher sensitivity is utilized for H₂ detection, while the high-absorption FP resonance enhances the photothermal catalytic effect, thereby improving the dynamic response speed of H₂ detection. We elucidate theoretically the H₂ sensing mechanism of TPP by comparing it with FP resonance, providing a novel design scheme for multilayer planar sensors. Furthermore, the dependence of TPP's sensitivity on Pd film thickness has also been thoroughly analyzed and discussed theoretically and experimentally. Due to the improved reproducibility demonstrated by the FOTPP tip with a thicker Pd film, we highlight its sensing perfor-

mance, including sensitivity, repeatability, and stability. Compared to traditional FP resonance sensing structures, the FOTPP H₂ sensing tip exhibits excellent H₂ detection performance and minimal cross-sensitivity due to its independence from Pd film deformation. Additionally, we provide a comprehensive theoretical and experimental investigation of the high-efficiency heating and catalytic effects of FP resonance photothermal effect, including their dependencies on resonance wavelength, laser power, and Pd film thickness. The results indicate that the FP resonance-enhanced photothermal effect significantly accelerates the response and recovery speeds of the FOTPP sensing tip and surpasses the photothermal assistance provided by non-resonance modes. This provides the necessary theoretical and experimental validation for the photothermal-assisted detection system.

Structural design and analysis of sensing mechanism

Figure 1(a) and 1(b) depict three-dimensional schematic diagram and cross-section of the proposed FOTPP sensing tip, which consists of both dielectric DBR and H₂-sensitive metal Pd. The DBR is composed of 5 pairs of alternately stacked Al₂O₃ and TiO₂ dielectric layers, designed to create photonic bandgaps. The thicknesses of the TiO₂ and Al₂O₃ layers satisfy the Bragg condition: $n_ih_i = \lambda_{\text{Bragg}}/4$ (i = 2, 3), where n_i and h_i represent the refractive index and thickness of TiO₂ and Al₂O₃, respectively, and λ_{Bragg} represents the Bragg wavelength. In this

design, the values of n_2 and n_3 are measured by the ellipsometer (M-2000 V), while the complex refractive indices of Pd and PdH_x are taken from the reported literature⁴⁷, as shown in Supplementary information Fig. S1. Besides, λ_{Bragg} is fixed at 400 nm with a 42 nm thick TiO₂ layer and a 60 nm thick Al₂O₃ layer, and the thickness of the top Pd film is denoted as h_1 . The preparation of FOTPP tips is detailed in the "Sample preparation" section. The cross-sectional scanning electron microscope (SEM) image on the right side of Fig. 1(b) demonstrates the uniformity of periodically stacked layers. Additionally, the metallic Pd not only excites TPP at the Pd/DBR interface but also serves as an H2 sensing layer. As shown in Fig. 1(c), there are both TPP and FP resonance modes generated in the FOTPP tip. The hydrogenation of the Pd film causes a redshift in the TPP resonance wavelength, thereby enabling H₂ detection through wavelength demodulation. Due to the inherent response recovery characteristics of Pd film, the FOTPP sensing tip also exhibits good repeatability. Furthermore, the photothermal effect induced by the FP resonance mode is employed to improve the response recovery speed of the FOTPP sensing tip.

Optical properties of the FOTPP tip are calculated by both the transfer matrix method (TMM) and the finitedifference time-domain (FDTD) method. Figure 2(a) shows the TMM-calculated dependence of reflection spectra on the thickness of Pd film. When there is no Pd film on the FOTPP tip, its reflection spectrum exhibits a



Fig. 1 | Photothermal synergistic TPP H₂ detection system integrated on the fiber tip. (**a**) Three-dimensional schematic of the fabricated fiber-optic TPP (FOTPP) tip with broadband halogen source (blue beam) and narrowband laser (red beam), along with a photograph of the actual fabricated FOTPP tip. (**b**) Its cross-section and SEM image for the FOTPP tip, where h_1 , h_2 , and h_3 denote the thicknesses of Pd, TiO₂, and Al₂O₃ layers, respectively. (**c**) Schematic of spectral working principle of photothermal synergistic FOTPP H₂ sensing tip and the response/recovery characteristics of resonance wavelength under photothermal (red curve) and non-photothermal (black curve) synergistic conditions.

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primary bandgap around 400 nm. With the increased thickness of the Pd film, a resonance dip (labeled as D_1) appears within this bandgap, along with two resonance dips at longer wavelengths labeled as D_2 and D_3 . The resonance depth of dip D_1 increases with the thickness of the Pd film, reaching a maximum near 50 nm. Besides, the effects of other structural parameters on the characteristic spectra are detailed in Supplementary information Fig. S2. For a direct comparison, the DBR is equivalent to a dielectric layer with an average RI of $\bar{n} = (n_2h_2 + n_3h_3)/(h_2 + h_3)^{48}$, serving as an FP resonance structure. Unlike the TPP resonance structure, the reflection spectra of the FP resonance structure exhibit four distinct orders of FP resonance dips, labeled as D_1' , D_2' , D_3' , and D_4' , as illustrated in Fig. 2(b). We further compared the H₂ detection sensitivity of the two different structures using TMM to demonstrate the superiority of the multilayer TPP resonance structure. Figure 2(c) illustrates the wavelength shift of the resonance dips of TPP (top panel) and FP (bottom panel) resonance struc-

tures after the hydrogenation of Pd films with different thicknesses. The results indicate that resonance dip D_1 exhibits higher sensitivity compared to dips D_2 and D_3 . Furthermore, the sensitivity of the resonance dip D_1 decreases rapidly as the Pd film thickness increases before stabilizing. Therefore, considering potential processing errors, it is crucial to use a sufficiently thick Pd film to ensure high reproducibility in the sensitivity of FOTPP tips. Furthermore, the sensitivity of the FP resonance structure is comparable to that of D_2 and D_3 , but less than half of the resonance dip D_1 mode in the FOTPP resonance structure for different thicknesses of Pd film. Therefore, the proposed FOTPP resonance structure demonstrates superior performance compared to the FP resonance structure for H₂ detection. The generation mechanisms of the resonance dips D_1 , D_2 , and D_3 are investigated through simulated electric field distribution. As shown in Fig. 2(d-f), resonance dip D_1 exhibits the strongest electric field at the DBR/Pd interface, gradually decaying within the DBR with a periodically modulated



Fig. 2 | Photoelectric characteristics of the FOTPP tip for H₂ detection. TMM calculated the reflection spectra of (**a**) FOTPP and (**b**) FP tips with different thicknesses of Pd film. The white dashed line marks the main bandgap location of the DBR. (**c**) Wavelength shifts of the resonance dips of TPP (top panel) and FP (bottom panel) tips after hydrogenation at various thicknesses of Pd film. (**d**–**f**) Simulated electric field profiles for three resonance dips supported by the FOTPP tip with a 50 nm thick Pd film, where E_0 denotes the incident electric field intensity.

exponential envelope, a typical characteristic of TPP. In contrast, resonance dips D_2 and D_3 produce interference fringes with highly uniform intensity, indicating their origin from different orders of FP resonance. Consequently, the sensitivity of the TPP mode surpasses that of the FP resonance mode.

To explain this phenomenon, the response mechanism of TPP to the hydrogenation of the Pd film is explored theoretically. Figure 3(a) depicts the electric field distribution curves of TPP before and after hydrogenation for a 50 nm thick Pd film. These results indicate that, after Pd hydrogenation, the electric field intensity of the TPP weakens, accompanied by a decrease in the penetration depth of the TPP into the DBR (L_{Bragg}). Here, the penetration depth is defined as the minimum of the electric field envelope. Additionally, Fig. 3(b, c) illustrates the dependence of the TPP electric field on the thickness of Pd film before and after hydrogenation, while Fig. 3(d) summarizes the maximum electric field intensity of TPP at the Pd/DBR interface. Both L_{Bragg} and

the electric field intensity of TPP mode increase with the thickness of the Pd/PdH_x layer, meanwhile they tend to stabilize as the thickness of Pd/PdH_x approaches 50 nm, indicating a correlation with reflection intensity. Additionally, both L_{Bragg} and the electric field intensity of TPP before hydrogenation are always greater than those after hydrogenation, which is attributed to the reduced reflection intensity of Pd film after hydrogenation, as shown in Fig. 3(e). Therefore, L_{Bragg} is positively correlated with the reflection intensity of the metal film on top of the DBR because greater reflection intensity excites stronger TPP resonance, in turn requiring a longer L_{Bragg} for attenuation. This phenomenon is key to achieving the H2induced wavelength shift of the TPP resonance. We further elucidate the response mechanism of TPP mode through its phase-matching condition as below⁴⁹:

$$2\bar{n}L_{
m Bragg}\cdot rac{\omega-\omega_{
m Bragg}}{c}=arphi_{
m Pd/PdH_x}\,,$$
 (1)

where ω and ω_{Bragg} is the resonance angular frequency of TPP and Bragg frequency, *c* is the speed of light, and



Fig. 3 | Response mechanisms of resonance wavelength in TPP mode. (a) Calculated electric field distribution at the resonance wavelength of TPP supported by the FOTPP tip before (top panel) and after hydrogenation (bottom panel). The dependence of the electric field distribution of TPP (b) before and (c) after hydrogenation on the thickness of the Pd film. The individual electric field curves are shifted upwards for clarity. The circle marks indicate the position range of L_{Bragg} (near the minimum of the electric field envelope). (d) The maximum electric field intensity of TPP at the Pd/DBR interface as a function of Pd film thickness. (e) Reflection intensity difference and (f) reflection phase difference between PdH_x and Pd films at different thicknesses. (g) Dependency of the total phase contribution of the DBR on the penetration depth of TPP in DBR (L_{Bragg}).

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$$2\bar{n}h\cdot\frac{\omega'}{c}+\varphi_{\mathrm{Pd/PdH}_{x}}=2\pi N\,,\qquad(2)$$

where ω' is the angular frequency of FP resonance mode, h is the total thickness of DBR, and N is an integer. Because the above two equations for both resonance modes incorporate the total phase contribution of the metal film, the reflection phases of the Pd and PdH_x films are calculated and shown in Fig. S3. Figure 3(f) illustrates the reflection phase difference ($\varphi_{PdHx} - \varphi_{Pd}$) before and after the hydrogenation of the Pd film. The results show that the change in the reflection phase of Pd films with different thicknesses before and after hydrogenation is minor. According to Eq. (2), the variation of the frequency of the FP resonance mode depends mainly on the difference between φ_{PdHx} and φ_{Pd} (other terms are constant or have negligible variation). Therefore, the wavelength shift of the FP resonance mode is relatively small. Different from the FP resonance mode, the wavelength shift of the TPP resonance mode mainly originates from the variation of L_{Bragg} . As depicted in Fig. 3(g), the L_{Bragg} variation significantly affects the total phase contribution of the DBR, resulting in a significant resonance wavelength shift of the TPP. Differentiating Eq. (1) with respect to L_{Bragg} , we have:

$$\frac{\mathrm{d}\omega}{\mathrm{d}L_{\mathrm{Bragg}}} = \frac{c}{2\bar{n}L_{\mathrm{Bragg}}} \left(-\frac{\varphi_{\mathrm{Pd/PdH}_x}}{L_{\mathrm{Bragg}}} + \frac{\mathrm{d}\varphi_{\mathrm{Pd/PdH}_x}}{\mathrm{d}L_{\mathrm{Bragg}}} \right), \quad (3)$$

where $d\varphi_{Pd/PdHx} / dL_{Bragg}$ approaches to 0 ($d\varphi_{Pd/PdHx}$ is minor) as well as both φ_{Pd} and φ_{PdHx} are always negative, this derivative is always greater than 0, that is, the resonance angular frequency of TPP is positively correlated with L_{Bragg} . Therefore, we demonstrate that the hydrogenation of the Pd film results in a reduction of L_{Bragg} , thus causing the redshift of TPP resonance wavelength. Moreover, the second-order differential of Eq. (1) is approximated as follows:

$$\frac{\mathrm{d}^{2}\omega}{\mathrm{d}L_{\mathrm{Bragg}^{2}}} = \frac{c}{2\bar{n}L_{\mathrm{Bragg}}}$$

$$\left(\frac{\mathrm{d}^{2}\varphi_{\mathrm{Pd/PdH}_{x}}}{\mathrm{d}L_{\mathrm{Bragg}}^{2}} - \frac{2\mathrm{d}\varphi_{\mathrm{Pd/PdH}_{x}}}{L_{\mathrm{Bragg}}\mathrm{d}L_{\mathrm{Bragg}}} + \frac{2\varphi_{\mathrm{Pd/PdH}_{x}}}{L_{\mathrm{Bragg}}^{2}}\right)$$

$$\approx \frac{c\varphi_{\mathrm{Pd/PdH}_{x}}}{\bar{n}} \cdot \frac{1}{L_{\mathrm{Bragg}}^{3}} \qquad . \tag{4}$$

Since Eq. (4) is always less than 0, the resonance frequency of TPP exhibits a greater variation rate when L_{Bragg} is smaller. As previously mentioned, L_{Bragg} is positively correlated with Pd film's thickness. This indicates that the sensitivity of TPP is negatively correlated with the thickness of the Pd layer when the Pd thickness is relatively small, which is consistent with the result of Fig. 2(c). However, in this case, L_{Bragg} has a strong dependence on the Pd film's thickness, resulting in poor robustness of the FOTPP tip's sensitivity. The strong dependence of sensitivity on the thickness of the H2-sensitive material is also a common issue for many H₂ sensors. To address this issue, the thickness of the Pd film is increased to 50 nm in the experiment to minimize the dependence of L_{Bragg} on Pd film thickness, ensuring that the fabricated FOTPP tip has relatively stable sensitivity within the permissible error range. Furthermore, as shown in Fig. 2(c), the FOTPP tip exhibits more than twice the sensitivity compared to the FP resonance structure, even for the thicker Pd film. Therefore, the FOTPP tip with a thicker Pd film simultaneously offers the advantages of high sensitivity and high reproducibility in fabrication.

Hydrogen sensing application of photothermal synergistic FOTPP tip

The fabricated FOTPP tip is employed in H₂ detection to experimentally validate the rationalization of the proposed theory. Figure 4(a) illustrates the schematic diagram of the detection setup. Different H₂ concentrations are prepared using the mixture of high-purity nitrogen and high-purity H₂ in various proportions in a gas chamber controlled by two mass flow meters. The FOTPP sensing tip is placed inside the gas chamber, and a terminal reflection spectroscopy system captures the spectra and dynamically collects the resonance wavelength of TPP. As shown in Fig. 4(b), the experimentally measured reflection spectra of the FOTPP tip (top panel), without a Pd film and with a 50 nm thick Pd film, exhibit spectral characteristics consistent with theoretical alculations (bottom panel). This indicates that the designed FOTPP tip can support three distinct resonance modes simultaneously. The resonance intensity discrepancy of dip D_1 is mainly attributed to the differences in the permittivity of the metal Pd, as well as thickness deviation, roughness, and imperfect uniformity of multilayer film. The sensitivity of the FOTPP tip is further assessed by placing it in different H_2 concentrations. Figure 4(c)

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illustrates the experimental spectra of the FOTPP tip at varying H₂ concentrations, revealing a redshift of TPP with increasing H₂ concentration. To demonstrate the response recovery properties and signal-to-noise ratio of the FOTPP tips, the gas in the chamber is cyclically switched between 100% N2 and a specific H2 concentration, maintaining each time interval of 300 s. The H₂ concentration gradually increases from 0% to 3.5%, then decreases from 3.5% back to 0%. This detection concentration range enables real-time quantitative monitoring of H₂ concentration in applications such as H₂ production and storage facilities, fuel cell systems, and the chemical industry, thereby addressing the safety requirements in industrial and energy storage and other fields³. Besides, according to our laboratory's safety management regulations, the case of H₂ concentration excessing 4% is not considered. Figure 4(d) and 4(e) show the amount of wavelength redshifts and real-time wavelength response of the FOTPP tip during the process. Experimental results indicate that the resonance wavelength

of TPP is approximately linearly correlated with the H₂ concentration, with the sensitivity S for both cycles remaining nearly consistent at approximately 0.661 nm/1%. The discrepancy between experimental and theoretical sensitivities is attributed to the incomplete hydrogenation of the Pd film and processing errors. In general, the limit of detection is defined as $3\sigma/S$, where σ represents the standard deviation of the baseline (~0.0163 nm, as shown in Supplementary information Fig. S5(b)). Therefore, the ideal limit of detection of the sensor is determined to be 0.074%, showing the potential of FOTPP tips for detecting low H₂ concentrations. Additionally, after responding at different H₂ concentrations, the resonance wavelengths of TPP return to the initial positions in high-purity N₂, demonstrating good response recovery characteristics. Furthermore, we demonstrate that the resonance wavelength shift of TPP has a linear relationship with the external temperature, as shown in Fig. 4(f) and 4(g). Since the FOTPP H₂ sensor does not rely on the deformation of a suspended thin film, it exhibits a



Fig. 4 | Experimental measurements of H_2 concentration using the fabricated FOTPP tip. (**a**) Schematic of the experimental setup for H_2 detection, highlighting the gas sensing and spectral analysis systems in the green and blue boxes, respectively. (**b**) Experimentally measured (top panel) and theoretically calculated (bottom panel) reflection spectra of the FOTPP tip without a Pd film and with a 50 nm thick Pd film. (**c**) Reflection spectra of the FOTPP tip under various H_2 concentrations ranging from 0.5% to 3.5%. (**d**) Wavelength redshifts and (**e**) real-time response of wavelength shift for the FOTPP tip at both increasing and decreasing H_2 concentration pulses, ranging from 0.5% to 3.5% and 3.5% to 0.5%. (**f**) Real-time wavelength shift response of TPP supported by the FOTPP tip under different working temperatures. (**g**) The relationship between the amount of wavelength shifts of TPP and working temperatures. (**h**) The wavelength response of the FOTPP tip in continuously repeated 0.5% H_2 concentration. The black dashed line represents the locations of the average wavelength shifts.

low temperature sensitivity of just 0.0072 nm/°C. This indicates that the FOTPP tip possesses relatively small cross-sensitivity, thereby enhancing its suitability for practical environmental monitoring applications. Furthermore, the FOTPP tip undergoes 8 cycles of response and recovery pulses in a 0.5% H₂ concentration to demonstrate its repeatability, as depicted in Fig. 4(h). The average response of the FOTPP tip in multiple detections of the 0.5% H₂ concentration has a wavelength shift of 0.365 nm with a standard deviation of 0.0238 nm. The small standard deviation confirms the FOTPP tip capability for repeated detection of low concentrations of H₂ and its long-term durability. In addition to the FOTPP tip with a 50 nm thick Pd film, the sensing performance of FOTPP tips with 28 nm and 36 nm Pd films is also shown in Fig. S4. Experimentally, the trend in sensitivity variation, as illustrated in Fig. S4(c), is consistent with the theoretical calculations in Fig. 2(c). Moreover, to demonstrate the advantages of the FOTPP tip, the quantitative comparison of sensing performances between our work and other recent works is listed in Table S1. The comparison results indicate that the proposed FOTPP tip exhibits excellent sensing performance without the need for complex fabrication processes.

In addition to H₂ detection sensitivity, the dynamic response speed of the sensor is also an important factor in early warning applications for H₂ leakage. According to the Arrhenius equation⁵⁰, the rate of the H-Pd reaction is positively correlated with temperature. Therefore, increasing the surface temperature of the H₂ sensors is an effective method to improve its response time. Building on the multi-resonance characteristics of FOTPP tips, we further employ the FP resonance mode within the same structure to enhance light absorption, enabling a localized and efficient photothermal effect at the fiber tip while ensuring no crosstalk between photothermal assistance and H₂ sensing. Before further discussing the sensing performance of the photothermal-assisted FOTPP sensing tip, we will first provide a detailed demonstration of the photothermal effect in the FOTPP structure. Figure 5(a) illustrates the simulated absorption spectrum of the FOTPP tip before and after hydrogenation. There are three distinct absorption peaks corresponding to TPP and two different orders of FP resonance modes. As a representative example, a 785 nm laser (MDL-E-785-30 mW) located at the absorption peak is employed, while a 980 nm laser (LSR980H-4W-FC) situated at the absorption dip is used for comparison. The reasons for selecting a broad absorption peak rather than a narrow absorption peak (~550 nm) for photothermal-assisted effects are summarized as two aspects: (1) a broad absorption peak exhibits excellent tolerance to deviation between the laser wavelength and the absorption peak; (2) the photothermal effect enhanced by a broad absorption peak demonstrates greater stability and reduced temperature cross-sensitivity when the ambient environment changes. The loaded power of these two lasers measured using the optical power meter (S130VC) is 7 mW, as shown in Fig. S5(a). Figure 5(b) illustrates the effect of introducing these two lasers on the resonance wavelength of TPP. Due to the change in the permittivity of the materials induced by the generated photothermal effect, the TPP resonance wavelength undergoes a redshift. Specifically, the redshift induced by the 785 nm laser is larger, indicating a higher generating temperature. Moreover, we demonstrate that the FOTPP tip exhibits excellent stability when exposed to air for a prolonged duration, either under laser illumination or non-laser illumination (as shown in Fig. S5(b-d)). To quantitatively describe the temperatures generated by these two lasers, comprehensive photothermal numerical calculations are performed. The scaled absorbed power Qabs can be expressed by the following equation⁵¹:

$$Q_{\rm abs} = \frac{\varepsilon_0 \omega \varepsilon'' |E|^2}{\int \operatorname{Re}\left(P_{\rm inc}\right) \cdot \mathrm{d}S_{\rm sim}} \cdot I_{\rm Laser} \cdot S_{\rm sim} , \qquad (5)$$

where ε_0 and ε'' represent the vacuum permittivity and the imaginary part of the permittivity of the materials, respectively. E is the electric field intensity in the materials. Pinc and Ssim are the Poynting vectors and the illuminated area (simulated region) of the incident light, respectively. ILaser is the power density of the laser, defined as the ratio of laser power to spot area. Since the spot diameter is equal to the fiber core diameter (400 μ m), I_{Laser} is determined to be 5.573×10^4 W/m². Based on Eq. (5), the scaled absorption power distributions of the FOTPP tip at wavelengths of 785 nm and 980 nm are calculated and illustrated in Fig. 5(c) and 5(d). The absorbed power is concentrated within the Pd film, with the FOTPP tip showing greater absorbed power for the 785 nm laser compared to the 980 nm laser. This indicates that resonance modes can more effectively absorb light energy, thereby enhancing light-to-heat energy conversion. According to the distribution of absorption power, the thermodynamic properties of the FOTPP resonance structure are further performed using the finite element

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Fig. 5 | Theoretical and experimental characterization of photothermal effect in the FOTPP tip with a 50 nm thick Pd film. (a) Simulated absorption spectra of the FOTPP tip before and after hydrogenation. (b) The effect of two 7 mW lasers with different wavelengths on the resonance wavelength of TPP. Calculated absorbed power and temperature distributions at wavelengths of (c, d) 785 nm and (e, f) 980 nm. Experimentally measured thermal images of the FOTPP tip under (g) 785 nm and (h) 980 nm laser illumination. (i–k) Thermal images of the FOTPP tip illuminated with three higher-power 985 nm lasers (9.5 mW, 10.5 mW, and 11.5 mW). The peak temperature of the fiber-optic tip and the ambient temperature are indicated in the thermal images. (I) Simulated and measured peak temperature at the FOTPP tip under different laser powers. For clarity, scenarios illuminated by 7 mW and higher power lasers are differentiated by enclosing them in red and blue boxes, respectively.

method. Figure 5(e) and 5(f) illustrate the temperature distribution of the FOTPP structure under these two laser illuminations. The temperature within the multilayer structure is nearly uniform in steady state, with optical heating from the 785 nm laser and the 980 nm laser resulting in temperatures of approximately 43.5 °C and 35.9 °C, respectively. In the experiment, thermal images of the FOTPP tip illuminated with these two lasers (785 nm and 980 nm) are captured directly using a high-resolution infrared thermal camera (FOTRIC 220 s), as depicted in Fig. 5(g) and 5(h). The peak temperature at the fiber tip and the ambient temperature are also indicated in these figures. The experimental results are in near agreement with numerical simulations, thereby validating the relationship between the photothermal effect and the resonance wavelength. Additionally, Fig. 5(i-l) provide theoretical and experimental evidence that the photothermal effect is proportional to laser power. This result also shows that the laser with non-resonance wavelengths requires higher power to achieve the same photothermal effect. Therefore, the resonance absorption effect of the structure can effectively enhance and regulate the photothermal effect.

The photothermal-synergistic sensing setup for rapid H_2 detection is developed based on the aforementioned multi-resonance enhancement scheme. As depicted in Fig. 6(a), the laser and broadband halogen light source are simultaneously introduced into the FOTPP tip via fiber-optic jumpers. Additionally, a filter is placed in front of the spectrometer to cut off reflected light with wavelengths longer than 750 nm, preventing potential damage from reflected lasers. Noteworthy, all tests are conducted in a thermally stable environment, where the laser output power remains stable, leading to minimal temperature variations on the sensor surface. In view of the extremely small temperature sensitivity of TPP, the wavelength shift induced by slight temperature variations is negligible, with only a corresponding adjustment

of the sensor's baseline at different operating temperatures. Figure 6(b) shows the real-time wavelength shift response of the TPP mode under photothermal assistance from both 785 nm (top panel) and 980 nm (bottom panel) lasers, which indicates that the FOTPP tip maintains excellent response and recovery characteristics and signal-to-noise ratio under photothermal assistance, as like as that under non-photothermal conditions. To demonstrate the advantages of photothermalassisted H₂ detection, Fig. 6(c) demonstrates the response and recovery curve of TPP at a 3.5% H₂ concentration under both non-photothermal and photothermal conditions. Without the photothermal synergy, the response and recovery times of H₂ detection are 122 s and 179 s, respectively. However, with the photothermal assistance of a 785 nm laser, the response and recovery times are reduced by approximately 85% and 53% (~19 s and 85 s) and are shorter than those achieved with a 980 nm laser (~25 s and 104 s). These results demonstrate that the photothermal catalytic effect significantly improves the response and recovery speeds of the FOTPP tip, with the effect being more pronounced under resonance-enhanced photothermal conditions. Figure 6(d) and 6(e) summarize the response and recovery times of the FOTPP tip at various H₂ concentrations under three

different conditions. The results at other H₂ concentrations further highlight the superiority of resonance-enhanced photothermal assistance in improving the dynamic response/recovery speed. Furthermore, the dynamic response and recovery speed of the photothermalassisted FOTPP tip exhibits a positive correlation with H₂ concentration, as the adsorption and diffusion rates of H₂ molecules are faster at higher concentrations. The influence of the photothermal effect on the wavelength shift of the TPP is also thoroughly explored, as illustrated in Fig. 6(f). It is noteworthy that the photothermal effect reduces wavelength shifts of the TPP mode, particularly in high-concentration H₂ environments. This is because the increase in temperature causes the Pd lattice to expand, facilitating the diffusion of H₂ into the surrounding environment, thereby reducing the binding ratio of H and Pd. Despite this reduction in wavelength shift, the sensitivity remains sufficient for H₂ detection application. Moreover, recent research has indicated that an increase in temperature is beneficial for H₂ detection in high relative humidity environments8. Therefore, the photothermal effect can also provide an effective strategy for overcoming environmental interference such as high relative humidity, further highlighting the importance of developing photothermal synergistic detection



Fig. 6 | Quantitatively assessment of the sensing performance of resonance-enhanced photothermal synergistic FOTPP tip with a 50 nm thick Pd film. (a) Schematic of the optical setup used for photothermal assistance. (b) Real-time wavelengths shift of TPP mode supported by the FOTPP tip under 785 nm laser illumination (top panel) and 980 nm laser illumination (bottom panel). (c) Response recovery curve of TPP at a 3.5% H₂ concentration under non-photothermal (blue curve), 785 nm-photothermal (red curve), and 980 nm-photothermal (green curve) conditions. Comparison of (d) response time, (e) recovery time, and (f) wavelength redshifts for the FOTPP tip at various H₂ concentrations under three different photothermal conditions.

This is an early view version and will be formally published in a coming issue with other articles in due time.

technology. Finally, we discuss the effect of Pd film's thickness on the photothermal synergistic H_2 detection system, as depicted in Fig. S6. This theoretical and experimental evidence provides valuable insights into optimizing the photothermal-assisted detection technology.

Conclusions and discussion

In conclusion, we have successfully designed and fabricated a multi-resonance enhanced photothermal synergistic FOTPP H₂ sensing tip. Through numerical simulations and theoretical analysis, we have demonstrated that the TPP response to H₂ is primarily attributed to changes in phase-matching conditions, driven by the decreased penetration depth within the PC. This distinctive operational mechanism enhances the sensitivity of TPP to more than twice that of the FP resonance mode. Experimentally, the sensitivity of the FOTPP H₂ sensing tips surpasses many fiber-optic H₂ sensors relying on Pd film deformation, with extremely low temperature cross-sensitivity. Furthermore, the relationship between TPP sensitivity and Pd film thickness has been confirmed both theoretically and experimentally.

In addition to the TPP resonance mode, we also achieve an efficient photothermal effect through the FP resonance mode in the FOTPP structure, significantly catalyzing the reaction between Pd and H. Theoretical and experimental results demonstrate that FP resonance absorption effectively improves the optical heating induced by the photothermal effects, thereby further reducing the sensor's response and recovery times. In particular, the response and recovery speeds of the FOTPP sensor are improved by 6.5 times and 2.1 times, respectively, at an H₂ concentration of 3.5%. In terms of fabrication, the FOTPP sensing tip offers unparalleled advantages over traditional sensors, typically requiring lithography or structural transfer. It can be fabricated simply through thin film deposition, enabling cost-effective batch production of the sensor. Thus, our work not only paves a new path for integrated TPP devices but also offers crucial theoretical reinforcement and experimental verification for photothermal-assisted sensing technology.

Notably, in certain circumstances, molecules such as H_2O , CO, and NO_x can strongly bind to the surface of H_2 -sensitive materials, effectively inhibiting the adsorption and dissociation of H_2 , which ultimately leads to sensor deactivation. Fortunately, previous studies have demonstrated that elevated temperatures can effectively mitigate this deactivation issue⁸. Therefore, although the present study only focuses on sensing applications un-

der conventional environmental conditions, the precisely controllable photothermal synergistic detection technique provides a potential strategy to overcome the competitive adsorption of H_2O and other molecules, thereby enhancing the environmental adaptability of hydrogen sensors.

Materials and methods

Numerical simulation approach:

In the TMM calculation, the transfer matrix of each layer can be determined by the following forms:

$$M_{i} = \begin{pmatrix} \cos\delta_{i} & -\frac{\mathrm{i}}{\eta_{i}}\sin\delta_{i} \\ -\mathrm{i}\eta_{i}\sin\delta_{i} & \cos\delta_{i} \end{pmatrix}, \quad (6)$$

where δ_i and η_i satisfy the following equation:

$$\delta_i = \frac{\omega}{c} n_i h_i, \quad \eta_i = \sqrt{\frac{\varepsilon_0}{\mu_0}} n_i . \tag{7}$$

Among them, n_i and h_i represent the refractive index and thickness of each layer. ε_0 and μ_0 are vacuum permittivity and vacuum magnetic permeability. The transmission behavior of incident light in multilayer dielectrics is expressed as the multiplication of the transfer matrix of each dielectric layer:

$$\begin{pmatrix} E_{1} \\ H_{1} \end{pmatrix} = \left(\prod_{i=1}^{N} M_{i}\right) \begin{pmatrix} E_{N+1} \\ H_{N+1} \end{pmatrix}$$
$$= \begin{pmatrix} A & B \\ C & D \end{pmatrix} \begin{pmatrix} E_{N+1} \\ H_{N+1} \end{pmatrix}, \quad (8)$$

where A, B, C, D are the elements of the total matrix of 2×2 . The reflection coefficient of the multilayer structure can be written as follows:

$$r = \frac{A\eta_0 + B\eta_0\eta_{N+1} - C - D\eta_{N+1}}{A\eta_0 + B\eta_0\eta_{N+1} + C + D\eta_{N+1}} .$$
(9)

In the FDTD simulation, a two-dimensional simulation is employed in the x-z plane, with periodic boundary conditions and perfectly matched layers applied along the x and z directions. The incident light has an electric field amplitude of 1 V/m, with the incident and polarization directions along the z and x axes, respectively.

In the thermal simulation, the temperature of the ambient surroundings is fixed to 24 °C. We assume that the temperature returns to the ambient temperature after propagating 1 mm inside the fiber core. Besides, the top of the FOTPP tip is cooled convectively via natural convection with a convective heat transfer coefficient of 10 $W/(m^2 \cdot K)$. The material parameters used in the thermal simulations are summarized in Table 1.

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Materials	Density (kg·m⁻₃)	Specific heat (J·(kg·K)⁻¹)	Thermal conductivity (W·(m·K) ⁻¹)
Pd	12023	240	72
TiO ₂	4950	690	7.4
Al ₂ O ₃	3900	796	28
SiO ₂	2203	709	1.38

Table 2	The coating	process	parameters	of the	atomic	layer	deposition	technique
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Thin film	Precursors	Pulse (ms)	Purge (s)	Precursor temperature (°C)	Deposition temperature (°C)	Cycle number
Al ₂ O ₃	Trimethylaluminum	20	20	Room temperature	150	580
	Water	20	25	Room temperature	150	
TiO ₂	Titanium tetrakis(dimethylamide)	100	20	70	150	715
	Water	20	25	Room temperature	100	

Material:

The multimode optical fiber for sensing is purchased from Changfei Optical Fiber & Cable Co., Ltd, with core/cladding/coating diameters of 400/430/730 μ m and a numerical aperture (NA) of 0.37. The materials of the core, cladding, and coating are high-purity quartz, fluorine-containing acrylic resin, and Ethylene-tetrafluoroethylene, respectively, and the optical fiber cutter was purchased from OSCOM Technology Co., Ltd.

Sample Preparation:

Initially, commercial fiber optics are cut using an optical fiber cutter to achieve a clean and flat fiber end facet. Subsequently, 5 pairs of Al_2O_3 and TiO_2 thin films are sequentially deposited on the fiber end facet using the atomic layer deposition (MNT-S100Oz-L3S1) technique with a vacuum degree of 20 Pa. The specific coating process parameters are shown in Table 2. Finally, a single layer of Pd film is deposited on top of the DBR using magnetron sputtering (Quorum Q300T+) with the sputtering current of 40 mA for a time of 100 s.

References

- Ould Amrouche S, Rekioua D, Rekioua T et al. Overview of energy storage in renewable energy systems. *Int J Hydrogen Energy* 41, 20914–20927 (2016).
- Yue ML, Lambert H, Pahon E et al. Hydrogen energy systems: a critical review of technologies, applications, trends and challenges. *Renew Sustain Energy Rev* 146, 111180 (2021).
- Sripriya, Meda US. Market study of hydrogen sensors and sensing systems. *ECS Trans* 107, 4489–4502 (2022).
- Tittl A, Mai P, Taubert R et al. Palladium-based plasmonic perfect absorber in the visible wavelength range and its application to hydrogen sensing. *Nano Lett* **11**, 4366–4369 (2011).
- Baldi A, Narayan TC, Koh AL et al. *In situ* detection of hydrogen-induced phase transitions in individual palladium nanocrystals. *Nat Mater* **13**, 1143–1148 (2014).
- Matuschek M, Singh DP, Jeong HH et al. Chiral plasmonic hydrogen sensors. *Small* 14, 1702990 (2018).

- Wen L, Sun ZW, Zheng QL et al. On-chip ultrasensitive and rapid hydrogen sensing based on plasmon-induced hot electron-molecule interaction. *Light Sci Appl* 12, 76 (2023).
- Tomeček D, Moberg HK, Nilsson S et al. Neural network enabled nanoplasmonic hydrogen sensors with 100 ppm limit of detection in humid air. *Nat Commun* 15, 1208 (2024).
- Korotcenkov G, Han SD, Stetter JR. Review of electrochemical hydrogen sensors. *Chem Rev* 109, 1402–1433 (2009).
- Liu Q, Yao JY, Wang YP et al. Temperature dependent response/recovery characteristics of Pd/Ni thin film based hydrogen sensor. Sens Actuators B Chem 290, 544–550 (2019).
- Caucheteur C, Guo T, Albert J. Review of plasmonic fiber optic biochemical sensors: improving the limit of detection. *Anal Bioanal Chem* **407**, 3883–3897 (2015).
- Jing JY, Liu K, Jiang JF et al. Highly sensitive and stable probe refractometer based on configurable plasmonic resonance with nano-modified fiber core. *Opto-Electron Adv* 6, 220072 (2023).
- Wang Q, Wang L. Lab-on-fiber: plasmonic nano-arrays for sensing. *Nanoscale* 12, 7485–7499 (2020).
- Xiong YF, Xu F. Multifunctional integration on optical fiber tips: challenges and opportunities. *Adv Photonics* 2, 064001 (2020).
- Zhao Y, Tong RJ, Xia F et al. Current status of optical fiber biosensor based on surface plasmon resonance. *Biosens Bioelectron* **142**, 111505 (2019).
- Liu HH, Hu DJJ, Sun QZ, Wei L, Li KW et al. Specialty optical fibers for advanced sensing applications. *Opto-Electron Sci* 2, 220025 (2023).
- Jiang BQ, Hou YG, Wu JX, Ma YX, Gan XT et al. In-fiber photoelectric device based on graphene-coated tilted fiber grating. *Opto-Electron Sci* 2, 230012 (2023).
- Yin SY, Guo Q, Liu SR et al. Three-dimensional multichannel waveguide grating filters. *Opto-Electron Sci* 3, 240003 (2024).
- Zhang YN, Liu YX, Shi BF et al. Lateral offset single-mode fiberbased Fabry–Perot interferometers with Vernier effect for hydrogen sensing. *Anal Chem* **95**, 872–880 (2023).
- Luo JX, Liu S, Chen PJ et al. Highly sensitive hydrogen sensor based on an optical driven nanofilm resonator. ACS Appl Mater Interfaces 14, 29357–29365 (2022).
- Wang CQ, Han ZW, Wang CX et al. Highly sensitive fiber grating hydrogen sensor based on hydrogen-doped Pt/WO₃. Sens Actuators B Chem 404, 135250 (2024).
- Ye Z, Ruan HB, Hu XY et al. TBAOH intercalated WO₃ for highperformance optical fiber hydrogen sensor. *Int J Hydrogen Energy* 47, 28204–28211 (2022).
- Cai SS, Liu F, Wang RL et al. Narrow bandwidth fiber-optic spectral combs for renewable hydrogen detection. Sci China Inf

Wei XR et al. Opto-Electron Sci 4, 240029 (2025)

https://doi.org/10.29026/oes.2025.240029

Sci 63, 222401 (2020).

- Gu FX, Wu GQ, Zeng HP. Hybrid photon–plasmon Mach–Zehnder interferometers for highly sensitive hydrogen sensing. *Nanoscale* 7, 924–929 (2015).
- Perrotton C, Westerwaal RJ, Javahiraly N et al. A reliable, sensitive and fast optical fiber hydrogen sensor based on surface plasmon resonance. *Opt Express* 21, 382–390 (2013).
- Nugroho FAA, Eklund R, Nilsson S et al. A fiber-optic nanoplasmonic hydrogen sensor *via* pattern-transfer of nanofabricated PdAu alloy nanostructures. *Nanoscale* **10**, 20533–20539 (2018).
- Yun S, Oyama ST. Correlations in palladium membranes for hydrogen separation: a review. *J Memb Sci* 375, 28–45 (2011).
- Adhikari S, Efremova MV, Spaeth P et al. Single-particle photothermal circular dichroism and photothermal magnetic circular dichroism microscopy. *Nano Lett* 24, 5093–5103 (2024).
- Cui XM, Ruan QF, Zhuo XL et al. Photothermal nanomaterials: a powerful light-to-heat converter. *Chem Rev* **123**, 6891–6952 (2023).
- Luo J, Wu QL, Zhou L et al. Plasmon-induced hot carrier dynamics and utilization. *Photonics Insights* 2, R08 (2023).
- Polley N, Sardar S, Werner P et al. Photothermomechanical nanopump: a flow-through plasmonic sensor at the fiber tip. ACS Nano 17, 1403–1413 (2023).
- Stewart JW, Nebabu T, Mikkelsen MH. Control of nanoscale heat generation with lithography-free metasurface absorbers. *Nano Lett* 22, 5151–5157 (2022).
- Wu HT, Chen PW, Zhan XD et al. Marriage of a dual-plasmonic interface and optical microfiber for NIR-II cancer theranostics. *Adv Mater* 36, 2310571 (2024).
- Ye Z, Li Z, Dai JX et al. Hydrogen sensing performance investigations with optical heating and sensing element surface modification. *Int J Hydrogen Energy* 46, 1411–1419 (2021).
- Zhang XP, Li XT, Zhang XH et al. Photothermal-assisted hydrogen permeation enhancement. *Sens Actuators B Chem* 365, 131935 (2022).
- Kaliteevski M, Iorsh I, Brand S et al. Tamm plasmon-polaritons: possible electromagnetic states at the interface of a metal and a dielectric Bragg mirror. *Phys Rev B* 76, 165415 (2007).
- Lundt N, Klembt S, Cherotchenko E et al. Room-temperature Tamm-plasmon exciton-polaritons with a WSe₂ monolayer. *Nat Commun* 7, 13328 (2016).
- Hu MY, Zhang Y, Jiang X et al. Double-bowl state in photonic Dirac nodal line semimetal. *Light Sci Appl* **10**, 170 (2021).
- Kar C, Jena S, Udupa DV et al. Tamm plasmon polariton in planar structures: a brief overview and applications. *Opt Laser Technol* **159**, 108928 (2023).
- Normani S, Bertolotti P, Bisio F et al. Tamm plasmon resonance as optical fingerprint of silver/bacteria interaction. ACS Appl Mater Interfaces 15, 27750–27758 (2023).
- Sreekanth KV, Perumal J, Dinish US et al. Tunable Tamm plasmon cavity as a scalable biosensing platform for surface enhanced resonance Raman spectroscopy. *Nat Commun* 14, 7085 (2023).
- He MZ, Nolen JR, Nordlander J et al. Coupled Tamm phonon and plasmon polaritons for designer planar multiresonance absorbers. *Adv Mater* 35, 2209909 (2023).

- Wang ZY, Ho YL, Cao T et al. High-Q and tailorable fano resonances in a one-dimensional metal-optical Tamm state structure: from a narrowband perfect absorber to a narrowband perfect reflector. *Adv Funct Mater* **31**, 2102183 (2021).
- Lu H, Shi SH, Li DK et al. Strong self-enhancement of optical nonlinearity in a topological insulator with generation of Tamm state. *Laser Photonics Rev* 17, 2300269 (2023).
- Ko JH, Seo DH, Jeong HH et al. Sub-1-volt electrically programmable optical modulator based on active Tamm plasmon. *Adv Mater* 36, 2310556 (2024).
- Zhang KH, Chen ZY, Li HJ et al. A dual-band hydrogen sensor based on Tamm plasmon polaritons. *Phys Chem Chem Phys* 25, 20697–20705 (2023).
- Palm KJ, Murray JB, Narayan TC et al. Dynamic optical properties of metal hydrides. ACS Photonics 5, 4677–4686 (2018).
- Brovelli LR, Keller U. Simple analytical expressions for the reflectivity and the penetration depth of a Bragg mirror between arbitrary media. *Opt Commun* **116**, 343–350 (1995).
- Adams M, Cemlyn B, Henning I et al. Model for confined Tamm plasmon devices. J Opt Soc Am B 36, 125–130 (2019).
- Peleg M, Normand MD, Corradini MG. The Arrhenius equation revisited. *Crit Rev Food Sci Nutr* 52, 830–851 (2012).
- Chen X, Chen YT, Yan M et al. Nanosecond photothermal effects in plasmonic nanostructures. ACS Nano 6, 2550–2557 (2012).

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Author contributions

Xinran Wei conducted the numerical simulations, performed experimental measurements, and prepared the initial draft. Xuhui Zhang helped with the experimental measurement. Rui Li, Haonan Wei, Yijin He and Lanlan Shen assisted in data and results analysis. Yuzhang Liang, Ting Xu, Yurui Fang, and Wei Peng helped with theoretical analysis and supervised the project. Yuzhang Liang and Ting Xu supervised the manuscript writing. All authors participated in the discussion and confirmed the final manuscript.

Competing interests

The authors declare no competing financial interests.

Supplementary information

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