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Plasmonic Metal - Polymer Hybrid Nanomaterials for Ultrafast Hydrogen Detection

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Abstract:

Hydrogen-air mixtures are highly flammable. Therefore, hydrogen sensors are of paramount importance for timely leak detection during hydrogen handling and in a future hydrogen economy. However, none of the existing solutions meets the stringent performance targets set by stakeholders. Furthermore, deactivation due to poisoning by *e.g.* carbon monoxide is a widely unsolved problem. Here we present a plasmonic metal-polymer hybrid nanomaterial concept that makes it possible to reduce the apparent activation energy of hydrogen sorption in plasmonic nanoparticles by engineering the particle-polymer interface, and that provides unprecedented deactivation resistance *via* a tailored tandem polymer membrane. In concert with an optimized volume/surface ratio of the signal transducer uniquely offered by nanoparticles, this enables subsecond sensor response times. Simultaneously, hydrogen sorption hysteresis is suppressed, sensor limit of detection is enhanced, and sensor operation in demanding chemical environments is enabled, without signs of long-term deactivation. In a wider perspective, our work opens the door to next-generation optical gas sensors with functionalities optimized by hybrid material engineering.

Main Text:

In a hydrogen economy,¹ where hydrogen gas is the clean and sustainable carrier of energy, hydrogen sensors will play a critical role due to hydrogen's wide flammability range in air. For safety reasons, any leaks in hydrogen energy storage systems, vehicles and appliances, as well as the entire hydrogen distribution infrastructure, must be detected immediately. Hence, hydrogen sensor performance targets specify, among other, a response time of 1.0 s at room temperature, across the concentration range from 0.1% to $10\%^2$ (see corresponding discussion in Supplementary Information). In the quest to meet these challenging targets, optical nanoplasmonic hydrogen sensors based on hydride-forming metal nanoparticles have been introduced.³⁻⁷ They are attractive because the optical signals generate no sparks and stem from absorption of hydrogen species into interstitial sites of the metal host, which renders such sensors intrinsically highly hydrogen-selective.³ Furthermore, their optical fingerprint is spectrally tunable,^{4,8,9} and they can be miniaturized down to the single nanoparticle level.^{6,10,11} In this field, like in many other hydrogen sensor platforms,^{12,13} Pd is the functional material of choice. This is due to its ability to dissociate hydrogen gas efficiently at ambient conditions, and its reversible phase transformation from metal to metal hydride at room temperature,^{3,14} which gives rise to a sizable optical contrast.¹⁵ However, among several other drawbacks, such as hysteretic behavior¹⁶ and response times falling short of the target value,^{13,17} the problem that hydrogen dissociation on Pd is effectively poisoned even by trace amounts of species like CO and NO2^{17,18} remains widely unsolved, in particular in combination with other critical sensor metrics (Tables S1-3). This is problematic since e.g. CO is omnipresent at ca. 0.2 ppm, and at even higher concentrations close to large point sources or in urban areas.¹⁹

The plasmonic metal-polymer optical hydrogen sensor platform presented here overcomes all these long-standing limitations by capitalizing on generic and synergistic effects that arise from combining PdAu alloy plasmonic nanoparticle signal transducers with tailored thin polymer membrane layers (**Fig. 1**a). As we show below, the stringent 1.0 s room temperature response time target can be met by tailoring the nanoparticle surface/volume ratio in concert with a reduction of the apparent activation energy by engineering the particle-polymer interface. Simultaneously, the hysteresis is suppressed, the sensor limit of detection is significantly enhanced and sensor operation in demanding chemical environments without any signs of deactivation, even after 4 months, is enabled.

Plasmonic metal-polymer optical hydrogen sensor platform

The nanoarchitecture of our hybrid sensor consists of a nanofabricated quasi-random array of hydride-forming and plasmonically active metal nanoparticles on a glass substrate coated with a thin polymeric film (**Fig. 1a**-b). Specifically, we chose a Pd₇₀Au₃₀ alloy, as well as a pure Pd control as detector. This alloy composition is optimal because at a lower Au fraction sensor accuracy is compromised by hysteresis^{20,21} (*i.e.* the readout is nonspecific as it depends on the hydrogen pressure history), while at a higher Au fraction the sensitivity is reduced.^{4,22} We nanofabricated the alloy and Pd nanostructures with average nanoparticle dimensions of 190 nm diameter and 25 nm height (the particles of this size are structurally very stable and exhibit a well manifested plasmon-related light-absorption peak) according to the Hole-Mask Colloidal Lithography method. Subsequent depositions of the metal constituents matching the targeted composition, are followed by high-temperature annealing to induce the alloy formation.²¹ This approach yields homogeneously alloyed polycrystalline disk-shaped nanoparticles^{4,21} with narrow size distribution (**Figs. 1**b and S1-4).

As the first polymer coating we chose polytetrafluoroethylene (PTFE) owing to its high chemical resistance and hydrophobicity. RF-sputtering of a 30 nm thin layer from a PTFE target yields a conformal coating of the entire nanoparticle array (**Figs. 1**c and S7-8). The optical response of the Pd₇₀Au₃₀ and Pd systems, measured in transmission configuration, is shown before and after applying the PTFE coating (**Fig. 1**d-e). The corresponding optical extinction spectra feature a localized surface plasmon resonance (LSPR) peak, which broadens and shifts to the red after PTFE deposition, due to the high refractive index of the coating.²³ Upon exposure to 1000 mbar of hydrogen, this peak further broadens and red-shifts owing to the change in both volume and permittivity of the metal nanoparticles, induced by hydrogen population of the interstitial lattice sites.¹⁵ This LSPR peak shift, which is a linear function of the hydrogen concentration inside the nanoparticles,²² then serves as the basis for our nanoparticle-polymer hybrid optical hydrogen sensors.

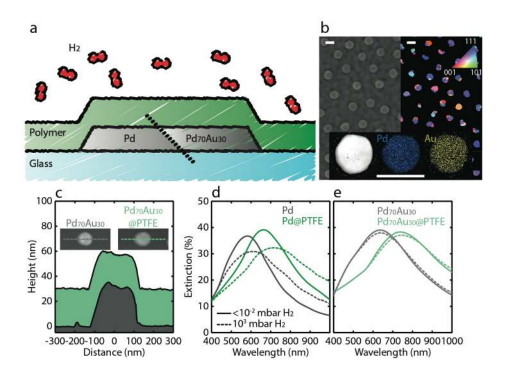


Figure 1. Plasmonic metal-polymer hybrid nanomaterial architecture and characterization. (a) Artist's rendition of a plasmonic metal-polymer nanomaterial comprising hydride-forming Pd or $Pd_{70}Au_{30}$ alloy nanoparticles, and a thin polymer coating. (b) SEM image (left) and Transmission Kikuchi Diffraction (TKD) micrographs (right) of a $Pd_{70}Au_{30}$ nanoparticle array, and EDX elemental maps of an individual $Pd_{70}Au_{30}$ alloy nanoparticle (bottom). The color code of the TKD images is explained in the inverse pole figure plot, which depicts the bulk grain crystallographic orientation with respect to the out-of-plane axis. Scale bars are 200 nm. (c) AFM image of an individual $Pd_{70}Au_{30}$ alloy particle before and after PTFE coating. The profile of the coated particle is shifted upward by the nominal 30 nm PTFE thickness for clarity. Optical extinction spectra of a (d) Pd and (e) $Pd_{70}Au_{30}$ alloy nanoparticle array before and after PTFE coating measured in vacuum and at 1000 mbar H₂ pressure.

Sensor@PTFE: accuracy and limit of detection

As the first characterization of our sensors, we measured optical hydrogen absorption and desorption isotherms at 30 °C for pure Pd and the Pd₇₀Au₃₀ alloy in the absence and presence of the 30 nm PTFE coating (**Fig. 2**a-b). For both systems the optical readout, *i.e.* the spectral shift of the plasmonic peak, $\Delta\lambda_{\text{peak}}$, is proportional to the H concentration in the host, expressed as

H/Pd ratio.^{15,22} For Pd, a characteristic α -phase region at low hydrogen partial pressure, where hydrogen is diluted at low concentration in a solid solution, is observed. At a critical hydrogen pressure, the $\alpha+\beta$ phase coexistence region ("plateau") at the first-order phase transition to and from the hydride (β -phase) appears and exhibits wide hysteresis. Finally, the pure β -phase region at high hydrogen partial pressure is observed.²⁴ In contrast, the Pd₇₀Au₃₀ system exhibits a monotonous hysteresis-free response. Interestingly, adding the PTFE coating to the two sensors not only retains the identified beneficial properties of the Pd₇₀Au₃₀ alloy but also enhances the signal amplitude by approximately a factor of two throughout the entire pressure range investigated. This is remarkable because, as becomes clear from quartz-crystal microbalance (QCM) measurements, this enhancement is not caused by hydrogen absorption in the PTFE itself (Fig. 2c-d). Specifically, our analysis indicates that the same amount of hydrogen is absorbed with and without PTFE coating both for Pd and the Pd₇₀Au₃₀ (the absolute numbers expressed as H/Pd ratio are in good agreement with earlier reports^{20,25}), corroborating that hydrogen only interacts with the nanoparticles. Furthermore, our Density Functional Theory (DFT) calculations show that the interaction between H₂ and PTFE (and PMMA) consists of only weak dispersion forces (Fig. S11).

This contrasts earlier work reporting on nearly doubled hydrogen solubility in MOF-coated Pd nanoparticles.²⁶ To address this aspect in more detail, we note that the sensitivity of a plasmonic hydrogen sensor is dictated by the spectral position of the plasmonic resonance, λ_{peak} , in the non-hydrogenated state.²² Since adding a polymer coating spectrally red-shifts λ_{peak} (cf. **Fig. 1**d-e), we expect an increase in sensitivity. This is indeed the case and the reason for the observed enhancement of the signal amplitude, as we discuss in detail in Supplementary Information based on Finite-Difference Time-Domain (FDTD) simulations. Since this enhancement is a purely optical far-field effect (the concentration of absorbed hydrogen remains the same), it is generic to *any* (polymer) coating and can be maximized by a high refractive index of the latter, as well as by increasing its thickness (Figs. S16-17) at the cost of a slight reduction of the figure-of-merit of the plasmonic sensor readout (Fig. S18). At the same time, it is exclusive to plasmonic hydrogen sensors based on nanoparticles since no such optical signal enhancement was observed for PTFE-coated Pd thin films.²⁷ Finally, we also note that the observed symmetric lowering of the plateau pressures for the Pd@PTFE system (cf. **Fig. 2**a and c) hints at strain imposed by the PTFE layer²⁸ (for the related estimates, see Supplementary Information)

The limit of detection (LoD, the lowest change in concentration of analyte that can be distinguished by a sensor) of the Pd@PTFE and Pd₇₀Au₃₀@PTFE sensors is derived by exposing them to pulses of gradually decreasing H₂ pressure in vacuum (from 1000 to 7 μ bar – the lowest pressure attainable in our setup) and by measuring the spectral shift of the plasmonic resonance peak maximum, $\Delta \lambda_{\text{peak}}$, at 1 Hz sampling frequency (Fig. 2e). Defining the LoD as 3σ , where σ is the noise of the acquired signal (i.e. 0.01 nm, see corresponding discussion in Supplementary Information and Fig. S20), we extrapolate that $LoD < 5 \mu bar$ in pure hydrogen (Fig. 2f). To assess the sensitivity under more realistic conditions, we also conducted similar experiments for the Pd₇₀Au₃₀@PTFE sensor in Ar and in synthetic air carrier gas at atmospheric pressure in a flow reactor setup. In both backgrounds, down to the lowest H₂ concentration experimentally attainable, *i.e.* 10 ppm and 20 ppm, respectively, a $\Delta \lambda_{peak}$ response is clearly discernible. This places our system amongst the most sensitive hydrogen sensors reported to-date under simulated application conditions in air,^{29,30} notably with significant potential to further enhance the sensitivity by e.g. tailoring the coating layer material and its thickness, and by optimizing nanoparticle dimensions.²² To this end, extrapolating the LoD based on the experimentally derived noise in flow conditions (Fig. S24), we derive an LoD of 1 ppm in Ar and 5 ppm in synthetic air.

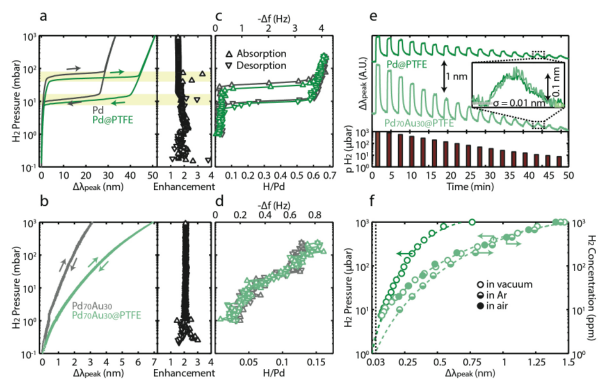


Figure 2. Sensitivity enhancement and limit of detection of Pd@PTFE and $Pd_{70}Au_{30}@PTFE$. Optical absorption and desorption isotherms of (a) Pd and (b) Pd₇₀Au₃₀ alloy sensors before and after PTFE coating. The arrows denote the sorption direction. The panels to the right depict the $\Delta \lambda_{peak}$ ratio of the coated and uncoated sensors, and reveal a factor of two enhanced signal amplitude induced by the PTFE. Isotherms of (c) Pd and (d) $Pd_{70}Au_{30}$ sensors before and after PTFE coating measured by quartz-crystal microbalance in a flow chamber at ambient pressure, using Ar as carrier gas. They reveal that the hydrogen concentration in the metal, expressed as *H*/*Pd*, is in excellent agreement with previous reports,²⁰ and that no H_2 is absorbed in the PTFE itself due to the identical response by the coated and uncoated system. (e) $\Delta \lambda_{peak}$ response to a step-wise decreasing H_2 pressure in the 7–1000 µbar range, measured at 1 Hz sampling frequency in a vacuum chamber. The inset shows the data at 10 µbar H_2 to define the noise, σ , as 0.01 nm (also see corresponding discussion in Supplementary Information). (f) Measured $\Delta \lambda_{peak}$ as a function of the H_2 pressure derived from (e) and also from measurements in Ar and synthetic air background down to the lowest attainable hydrogen partial pressure in our system, 10 ppm and 20 ppm, respectively (Figs. S25-26). The green lines depict an extrapolation from the lowest reliably attainable data point in our system for the Ar and air background to the 3σ point, indicating a LoD < 1 ppm and 5 ppm, respectively. The black dashed line marks the extrapolated LoD at $3\sigma = 0.03$ nm. All experiments were done at 30 °C.

Sensor@PTFE: response time

As the next characterization step, we measured the response times of uncoated and PTFE-coated Pd and Pd₇₀Au₃₀ alloy sensors at 30 °C, by monitoring their temporal response to a step-wise in/decrease of hydrogen pressure to/from 40 mbar H₂ (*i.e.* the lower flammability limit) in a vacuum chamber (**Figs. 3**a-b and S28 for recovery times). We observe two key effects, namely that (i) for both ab- and desorption the response time of the uncoated Pd₇₀Au₃₀ sensor is significantly shorter than for the uncoated Pd one, and (ii) the PTFE coating for both systems further shortens the response time to significantly below one second for the alloy. The detailed quantitative analysis of the Pd and Pd₇₀Au₃₀ ab- and desorption kinetics *via* Arrhenius analysis (Figs. S29-30) reveals a significant reduction of the apparent activation energies, *E_a*, of the rate-limiting steps for both hydrogen absorption and desorption. This is due to the PTFE coating

(**Figs. 3**d-e and S29-30), which also refines the understanding of the observed hydrogen sorption kinetics acceleration in PTFE-coated Pd thin films reported by Ngene *et al.*²⁷ These results, along with similar kinetics acceleration observed for MOF-coated Pd sensors measured in vacuum/hydrogen²⁶ and air,³¹ imply that the kinetics-accelerating effect of such coatings may be generic. To this end, a number of different explanations to the origin of the accelerated kinetics is given in the literature, such as the modification of surface chemical^{27,32} and electronic states^{26,27,33}, physical force/stress imposed by the coating layer³³ and the removal of competing molecules reacting on the surface.³¹

Employing DFT calculations to capture the experimentally observed trends, we show that the measured decrease in E_a induced by the polymer coating is connected to the absorption and desorption processes at the nanoparticle-polymer interface (**Fig. 3**e), mediated by polymer-metal bond formation verified by XPS analysis (Fig. S9). On one hand, the activation barrier for hydrogen absorption from surface to subsurface sites is reduced by 11 kJ/mol upon coating by PTFE, resulting in faster absorption. On the other hand, for sites close to PTFE, surface-adsorbed hydrogen is destabilized by 9 kJ/mol, which results in faster H₂ desorption during dehydrogenation. In other words, these effects give rise to a reduction of E_a for both hydrogen absorption and desorption barriers for the respective rate-limiting step. Concerning the kinetics, we also note that the dependence of the response time on pressure (**Fig. 3**g and **Figs. 4**e and **5**c below) is fairly weak and close to the power-law form with a small exponent (cf. Eq. S15). This kinetic feature is indicative of energetic heterogeneity at the surface of the alloy (see corresponding discussion in Supplementary Information).

Having established that the hydrogen absorption rate into the bulk of the nanoparticle sensors is limited by their surface,³⁴ the time scale for this process is expected to be proportional to their volume/surface ratio, with the proportionality constant depending on the specifics of the kinetics at the surface (Figs. S35 and S37 and the corresponding discussion). Hence, as a key advantage enabled by a sensor platform based on nanoparticles, tailoring the volume/surface ratio should render a faster response. Indeed, reducing the mean diameter of the nanoparticles in a $Pd_{70}Au_{30}@PTFE$ sensor from 190 nm to 100 nm further reduces the response time for desorption from ca. 9 s to 3 s (Fig. S28), and for absorption to (below) the resolution limit of our system at 0.3 s (Figs. 3c). Hence, the 100×25 nm $Pd_{70}Au_{30}@PTFE$ sensor meets the toughest of the US

DoE targets,² defined as response time of < 1 s to 1 mbar hydrogen at ambient temperature (**Fig. 3**f-g). Furthermore, a recovery time of below 5 s throughout the same hydrogen pressure range is also achieved (Fig. S32), representing the current state-of-the-art in this respect. To this end, we highlight that additional downsizing and volume/surface engineering of the nanostructures offers potential to further improve these already very impressive response times, in particular when using optimized nanoparticle designs, such as nanorods that combine a spectrally red-shifted LSPR for optimized LoD with small volume/surface, are used. However, on the note of utilizing the PTFE coating as selective membrane to prevent sensor deactivation by trace gases like CO and NO₂ present in air, we find that no sufficient protection is obtained. This is evidenced by the deactivation tests performed in a flow reactor operating at atmospheric pressure, using synthetic air as the carrier gas to mimic real application conditions. In particular, the sensor was exposed to 10 cycles of 4% H₂ followed by 10 cycles of 4% H₂ plus an interfering gas, *i.e.* 3% CO₂, 0.5% CH₄, 0.1% CO, 0.05% NO₂ (Fig. S49).

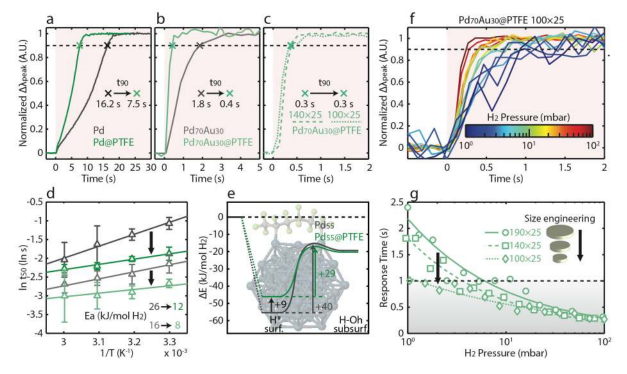


Figure 3. Response times of PTFE-coated Pd and $Pd_{70}Au_{30}$ at room temperature. Accelerated absorption kinetics are observed for PTFE-coated (a) Pd and (b) $Pd_{70}Au_{30}$ sensors with 190×25 nm average nanoparticle dimensions (first and second number denotes diameter and height, respectively). t_{90} is the response time defined as the time to reach 90% of the signal and is marked by black dashed lines. (c) Engineering the volume/surface ratio by employing

nanoparticles with smaller dimensions enables even faster absorption kinetics, as shown for 140×25 and 100×25 Pd₇₀Au₃₀@PTFE sensors. The shaded areas in panels (a-c) mark the period when the sensors are exposed to 40 mbar H₂. (d) Experimentally measured apparent hydrogen absorption activation energies, E_a , of a Pd and Pd₇₀Au₃₀ sensor before and after PTFE coating, revealing a distinct reduction of E_a upon PTFE coating. The error bars are the standard deviation from three measurements. (e) Energy landscape for hydrogen absorption in Pd₅₅ with and without PTFE adsorbed on the surface calculated by DFT. The figure depicts the states of adsorbed H on the surface (H* surf.) and absorbed H in a subsurface octahedral site (H-Oh subsurf.). The reference is taken as Pd₅₅+1/2H₂ (g) and Pd₅₅@PTFE+1/2H₂ (g), respectively. The background image depicts the modelled Pd₅₅@PTFE system. (f) 100×25 nm Pd₇₀Au₃₀@PTFE sensor response to H₂ pressure pulses in the range from 1000 to 1 mbar. (g) Response time of Pd₇₀Au₃₀@PTFE sensors with different sizes to varying H₂ pressure pulses. For the 100×25 nm sensor, sub-second response is achieved throughout the investigated pressure range down to 1 mbar, making this the fastest hydrogen sensor ever reported at these conditions (Table S2). The lines denote a power-law fit based on Eq. (S18).

Pd₇₀Au₃₀@PMMA Sensor

As a response to the shortcomings of the PTFE coating, in the second part of our study we investigate an alternative polymer system – poly(methyl methacrylate), PMMA, for which excellent H₂ selectivity to other gases has been demonstrated.^{35–37} To assess the molecular sieving function of a PMMA coating, we spin coated a 35 nm thick PMMA film onto a 190×25 Pd₇₀Au₃₀ sensor (for a neat Pd@PMMA analogue, see Supplementary Information). The corresponding Pd₇₀Au₃₀@PMMA sensor response in an identical deactivation test as described above for the PTFE sensor is summarized in **Fig. 4**a, revealing the excellent protection provided by the PMMA layer. Specifically, as depicted in **Fig. 4**b, the sensor signal for all interfering gases is with significant margin retained within the ±20% deviation limit from the normalized $\Delta\lambda_{peak}$ in pure 4% H₂ according to the performance standard for hydrogen sensors.³⁸ This is remarkable in view of the strong deactivation of the surface via strong CO chemisorption and the concurrent blocking of hydrogen dissociation sites.¹⁸

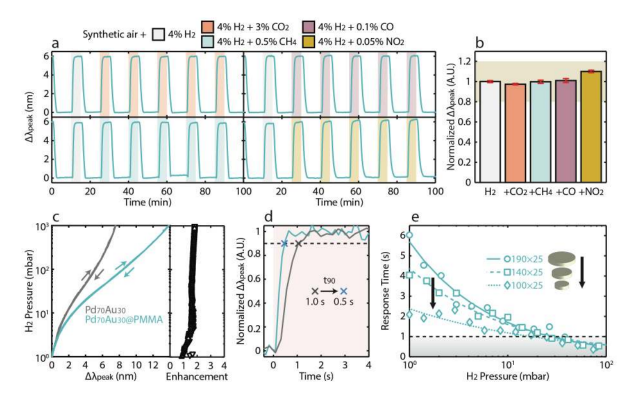


Figure 4. $Pd_{70}Au_{30}@PMMA Sensor. (a) Time-resolved <math>\Delta \lambda_{peak}$ response of $Pd_{70}Au_{30}@PMMA$ to a pulse of 4% H₂ followed by 5 pulses of, respectively, 4% H₂ and 3% CO₂, 4% H₂ and 0.5% CH₄, 4% H₂ and 0.1% CO, and 4% H₂ and 0.05% NO₂ in synthetic air carrier gas, measured at atmospheric pressure. The $\Delta \lambda_{peak}$ signals are retained throughout exposure to the different background gases, as summarized in (b) where the respective signals are normalized to the one obtained in pure 4% H₂. The error bars denote the standard deviation from 10 cycles. The shaded area indicates the ±20% deviation limit from the normalized $\Delta \lambda_{peak}$ in pure 4% H₂. (c) Optical absorption and desorption isotherms of the $Pd_{70}Au_{30}$ sensor measured before and after adding of the PMMA coating. The arrows denote the sorption direction. The panel to the right depicts the $\Delta \lambda_{peak}$ ratio of the coated and uncoated sensors, revealing a similar signal enhancement factor to the one found for the PTFE-coated system. (d) $Pd_{70}Au_{30}$ sensor response before and after PMMA coating to a step-wise H₂ pressure in/decrease to/from 40 mbar. (e) Response times of $Pd_{70}Au_{30}@PMMA$ sensors with different nanoparticle sizes to different H₂ pressure pulses. The lines denote a power-law fit based on Eq. (S18). All experiments were done at 30 °C.

Now, referring to the beneficial effects of the PTFE coating on the sensor performance identified above, we find that the same signal amplitude enhancement by approximately a factor of two is also observed for PMMA (Fig. 4c). Hence, as anticipated and discussed above, this is indeed a generic effect inherent to dielectric (polymer) coatings. The observed similar magnitude of the enhancement for PTFE and PMMA is thus solely the consequence of their similar refractive indices and the essentially identical thickness of the coatings, since also for PMMA no relevant hydrogen sorption occurs within the polymer itself (Fig. S11). In terms of response time shortening, our analysis reveals that this effect is also present for a PMMA coating, as the response time at 30 °C to a step-wise increase of the hydrogen pressure to 40 mbar H₂ for the Pd₇₀Au₃₀@PMMA sensor is reduced to 0.5 s (Fig. 4d). However, as more detailed analysis reveals, the acceleration of the hydrogen sorption kinetics is on average a factor two smaller for PMMA compared to PTFE and thus, even by reducing the nanoparticle diameter to 100 nm, the system falls slightly short of the DoE target for sub-second response to 1 mbar at room temperature (Fig. 4e). The reason we find is the smaller reduction of the apparent activation energy of the rate-limiting steps during hydrogen absorption and desorption induced by PMMA, compared to PTFE (Figs. S42-43), which is also confirmed by DFT calculations (Fig. S15).

Pd₇₀Au₃₀@PTFE/PMMA Tandem Sensor

As an intermediate conclusion it becomes clear that the ideal polymer layer essentially would *combine* the properties of the tested PTFE and PMMA systems. Hence, we fabricated a $Pd_{70}Au_{30}$ hydrogen sensor encapsulated by a *tandem* 30 nm PTFE + 35 nm PMMA structure. In this way, we are able to test the hypotheses that (i) it indeed is the PTFE-nanoparticle interface and the corresponding reduction of the apparent activation barriers that gives rise to the superior response time, (ii) PMMA functions as a superior molecular sieve layer, and (iii) these two functions can be combined in a heterostructure. To prepare the sensor, we applied a 5 s H₂-plasma treatment to the previously already tested $Pd_{70}Au_{30}$ @PTFE sensor (cf. **Figs. 1-3**) to render the PTFE-surface hydrophilic.³⁹ This enables wetting and thus spincoating of PMMA dissolved in methoxybenzene solvent. The obtained polymer tandem structure is schematically depicted in **Fig. 5**a, together with a cross-section SEM micrograph in which a PdAu alloy nanoparticle encapsulated in the tandem polymer arrangement is clearly resolved. The effective coating of the PTFE layer by the PMMA is further confirmed by XPS analysis (**Fig. 5**b).

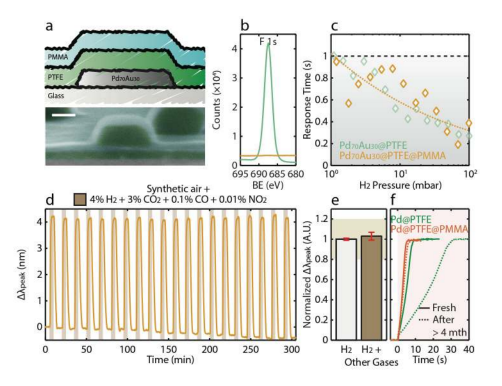


Figure 5. Pd₇₀Au₃₀@PTFE@PMMA Tandem Sensor. (a) Top: Artist's rendition of a tandem sensor comprising a heterostructure coating of PTFE and PMMA layers on top of a Pd₇₀Au₃₀ alloy nanoparticle array. Bottom: Cross-section SEM image of the tandem sensor revealing the polymer layers with different contrast. Scale bar is 100 nm. (b) XPS spectra of the F 1s peak of a $Pd_{70}Au_{30}(a)$ PTFE and a $Pd_{70}Au_{30}(a)$ PTFE(a) PMMA tandem sensor. The absence of the F 1s peak for the tandem confirms the effective coating of the PTFE by PMMA. (c) Response times of a tandem sensor with 100×25 nm nanoparticle dimensions to different H₂ pressure pulses, plotted together with the ones of the same sensor with only the PTFE coating. The dashed-line denotes a power-law fit based on Eq. (S18). (d) $\Delta\lambda_{peak}$ of the tandem sensor for 20 cycles of 4% H₂ plus 3% CO₂, 0.1% CO and 0.05% NO₂ in synthetic air. (e) $\Delta \lambda_{peak}$ in poisoning gas background normalized to $\Delta \lambda_{peak}$ in pure 4% H_2 revealing the excellent deactivation resistance of the tandem. The error bars denote the standard deviation from 20 cycles. The shaded area indicates the $\pm 20\%$ deviation limit from the normalized $\Delta \lambda_{peak}$ in pure 4% H₂. (f) Absorption kinetics of Pd@PTFE and Pd@PTFE@PMMA sensors to 40 mbar H_2 at different aging states. After four months in ambient conditions, the tandem sensor retains its kinetic properties while the PTFEonly-coated sensor has significantly deteriorated.

Assessing the room temperature response time in the same way as above reveals that for the tandem coating it is essentially identical to the PTFE coating alone (Fig. 5c). Since we use the same sensor in both cases also the reproducibility as such is remarkable. Furthermore, and as the key result, the sub-second response is again retained down to 1 mbar, meeting the DoE target. Comparing the apparent activation energies for the two systems, we find them to be identical (Fig. S46), corroborating that the kinetics acceleration observed indeed is governed solely by the direct interaction between coating material and nanoparticle surface. At the same time, the deactivation tests in synthetic air, following the same experimental scheme used above, reveal that the same deactivation resistance as for the PMMA alone is achieved (Fig. S49), convincingly confirming that the hybrid material concept is able to combine multiple functionalities of its constituents in a single system. Moreover, in a significantly exaggerated test, in terms of concentrations to "mimic" the environment of an urban area, the sensor with the tandem coating shows no sign of deactivation when exposed to twenty 4% H₂ pulses mixed with 3% CO₂, 0.1% CO and 0.01% NO₂ (Fig. 5d). Also in this tough test, the absolute response remains well within the \pm 20% deviation limit according to the performance standard for hydrogen sensors³⁸ (Fig. 5e). This excellent deactivation resistance is also retained in the long term. In particular, after four-month exposure to ambient conditions, the tandem sensor retains its response, while the only PTFE-coated sensor is significantly deactivated as indicated by drastically reduced response time (Fig. 5f). Finally, as an additional advantage of the tandem system, owing to its doubled polymer layer thickness, the sensitivity is further enhanced by 30% compared to the PTFE layer alone (Figs. S47-48), in agreement with our FDTD simulations (Fig. S17).

In summary, the concerted hydrogen sorption rate and sensitivity enhancement, as well as molecular sieving effects, we find for polymer coating layers on hydride-forming metal nanoparticles provide a new mechanism by which plasmonic nanoparticle arrays can operate as optical hydrogen sensors with unprecedented response metrics. The performance of the plasmonic metal-polymer optical hydrogen sensors presented here challenges existing hydrogen-sensing technologies (Table S2). Specifically, they (i) feature sub 5-ppm detection limit, (ii) exhibit deactivation resistance by poisoning gases such as O₂, CO₂, CH₄, NO₂ and CO, (iii) feature remote optical readout without risk for spark generation, (iv) are hysteresis-free, (v) exhibit excellent long-term stability, and (vi) meet for the first time the stringent 1 s response

time target at ambient temperature and 1 mbar H₂ partial pressure. The last point is achieved by concerting two key effects: (i) reducing the activation barrier for surface-to-subsurface hydrogen diffusion via polymer-metal surface bond formation; and (ii) tailoring the volume/surface ratio, a feature uniquely possible in nanoparticles, to overcome the intrinsic response time limitations imposed by the identified surface-associated rate-limiting steps during (de)hydrogenation of the sensor. The latter point also implies that it is fundamentally impossible to reach the 1 s response time target at low hydrogen partial pressures with macroscopic (bulk) or, to some extent, thin-film systems (see corresponding discussion in Supplementary Information) based on Pd and its alloys, since in this regime the volume/surface ratio is the critical factor determining the response time. In a wider perspective, our work opens the door to next-generation (optical) gas sensors centered on the idea of a hybrid material that combines tailored plasmonic signal transducers with multiple selective membrane materials which, by engineering the dimensions and interfaces of the material constituents, enable the optimization of sensitivity, selectivity, deactivation resistance and response times, to meet the ever increasing demand for advanced sensor technologies.

Methods

Sensor fabrication. Evaporation masks were fabricated using the standard hole-mask colloidal Lithography (HCL) process⁴⁰ on 1×1 cm² glass substrates (Borofloat, Schott Scandinavia AB), on TEM "windows" made in-house following the procedure by Grant *et al.*,⁴¹ on silicon wafer substrates and on QCM crystals (Laptech, SC-cut, 10 MHz fundamental frequency) pre-coated with 100 nm CVD-grown SiO₂ (PECVD, STS) on one of the Au electrodes. The steps of the mask fabrication were as follows: (i) substrates were cleaned (by ultrasonic agitation for the case of glass, silicon and QCM crystals and only by rinsing for the TEM windows in order to not break the membrane) subsequently in acetone, isopropanol (IPA), and de-ionized (DI) water. (ii) Poly(methyl methacrylate) (PMMA, MicroChem Corporation, 4 wt. % diluted in anisole, $M_W =$ 950 000) was spin-coated onto the substrates at 2000 rpm for 30 s (yielding a PMMA thickness of ~280 nm) followed by a soft baking at 170 °C on a hotplate for 5 min. (iii) Samples were subjected to a 5 s oxygen plasma (50 W, 250 mTorr, Plasma-Therm Batchtop RIE 95m) to enhance the hydrophilicity of the sample surface. (iv) A polyelectrolyte solution (poly diallyldimethylammonium (PDDA) $M_W = 200\ 000 \div 350\ 000$, Sigma Aldrich, 0.2 wt. % in Milli-Q water, Millipore) was pipetted on the surface of the samples and left incubating for 40 s before rinsing in DI water. This created a positively charged surface layer on the PMMA surface. (v) A suspension of negatively charged polystyrene beads (PS 190 nm sulfate latex, Interfacial Dynamics Corporation, 0.2 wt. % in Milli-Q water, Millipore) was added to the surface. The size of the PS beads determines the diameter of the fabricated nanodisks at the end of the processing. After 3 min incubation the suspension was rinsed away using DI water, and the samples were blown dry with nitrogen gas. (vi) A 15 nm thick Cr film was evaporated using a Lesker PVD 225

Evaporator at a base pressure of 5×10^{-7} Torr, evaporation rate 1 Å s⁻¹. (vii) The PS beads were removed by tape stripping (SWT-10, Nitto Scandinavia AB) for glass, silicon and QCM samples and by a wet tissue for TEM windows. This left a Cr film with holes at the positions of the stripped PS beads. (viii) The samples were subjected to 5 min oxygen plasma treatment (50 W, 250 mTorr, Plasma-Therm Batchtop RIE 95m) to etch through the PMMA layer exposed beneath the holes in the Cr mask. (ix) Through this mask Pd was deposited at a deposition rate of 1 Å s⁻¹. To fabricate the PdAu alloys, Au and Pd were deposited in sequence at the same deposition rate through the mask with tailored amount. The thicknesses of each Au and Pd layer determined the final composition of the alloy particles.²¹ (x) The remaining PMMA layer was dissolved in acetone in a lift-off step, removing the mask from the sample, only leaving the nanodisk structures on the substrate. (xi) Samples were soaked in IPA and blown dry with nitrogen. (xii) Both Pd and Au-Pd samples were annealed in a homemade flow furnace under 4% H₂ in Ar (100 ml min⁻¹) at 500 °C for 24 h, except for samples on QCM crystals, which were annealed instead at 400 °C for 72 h to avoid α - β phase transformation in quartz at 450 °C that leads to cracking. This way alloying of PdAu samples was achieved.

PTFE deposition. Thin PTFE films were prepared at room temperature in an ultrahigh-vacuum (UHV) DC/RF magnetron sputtering system with a base pressure of 10^{-7} mbar and a deposition pressure of 5×10^{-3} mbar Ar.⁴² The deposition rate was first determined by sputtering the target independently at a fixed power over a well-defined time interval. The thickness of the reference film was then measured with a DekTak3 profilometer, and the deposition rate (nm s⁻¹) was calculated from the thickness and time.

PMMA deposition. PMMA (MicroChem Corporation, 1 wt. % diluted in anisole, $M_W = 950$ 000) was spincoated on sensors at 2000 rpm for 30 s followed by a soft baking at 170 °C on a hotplate for 5 min resulted in 35 nm PMMA film, as measured by ellipsometry (J.A. Woollam M2000). To enable spincoating on an already PTFE-coated sensor to create the tandem structure, a hydrogen plasma treatment was applied for 5 s (100 W, 250 mTorr, Plasma-Therm Batchtop RIE 95m). This step rendered the PTFE surface hydrophilic and thus allowed spincoating of PMMA onto it. The PMMA deposition steps followed the ones described above.

Material characterization. Scanning Electron Microscopy (SEM): All SEM images were obtained in a Zeiss Supra 60 VP with secondary electron detector, working distance 4 mm, and an electron beam acceleration voltage of 10-15 kV. Scanning Transmission Electron Microscopy (STEM) and Energy Dispersive X-ray Spectroscopy (EDS): STEM-EDS measurements were taken using an FEI Titan TEM instrument operated at 300 kV acceleration voltage. The electron probe diameter and convergence semi-angle were approximately 0.15 nm and 17.5 mrad, respectively. HAADF-STEM images were recorded with a collection semi-angle of approximately 47 mrad. EDS maps were obtained with a pixel size of ~5 nm and acquisition time of 1 s per pixel. The EDS maps were analyzed in Aztec 3.3 (Oxford Instruments). Transmission Kikuchi Diffraction (TKD): The TKD investigation of the samples was performed on an FEI Nova Nano Lab 600 scanning electron microscope (SEM) equipped with Bruker OPTIMUSTM TKD detector operated at an acceleration voltage of 30 kV and beam current of 1.7 nA using an aperture of 30 µm. The electron transparent TEM window containing the nanoparticles was positioned horizontally on the microscope with the particles facing downward, and TKD orientation maps were acquired over a large number of particles with a scan step of 5 nm, and exposure time of approximately 9 ms per point with a pattern resolution of 320×240 pixels (5×5 binning). X-rav Photoelectron Spectroscopy (XPS): High-resolution (HR) XPS

spectra were recorded with a PHI 5000 system (Physical electronics, Chanhassen, Ma, USA) using monochromatized Al Ka radiation (hv1/4 1486.6 eV) as the x-ray source. The HR spectra displayed here were obtained at an incident angle of 45° with a combined energy resolution of 0.01 eV. Atomic sensitivity factors for quantitative analysis were taken from a reference.⁴³ The binding energies were corrected according to the Au 4f peak (84 eV) and the C 1s peak (284.8 eV). The Au 4f peak was also employed to correct for any additional charge build during HR XPS measurements. For all measurements, the samples were electrically grounded to prevent accumulation of electrostatic charge. Atomic Force Microscopy (AFM): AFM measurements were performed in tapping mode using an SPM Bruker Dimension 3100 system in air. In situ Xray Diffraction (XRD): The in situ XRD measurements were performed with a Bruker D8 Advance (Co-K α λ = 0.1789 nm) that is equipped with an LYNXEYE 1D detector. During the XRD measurements, the sample was hydrogenated at a constant temperature of 25 °C inside an Anton Paar XRK 900 reactor chamber. As the loading gas, a mixture of 96.0% helium and 4.0% H₂ was used and a constant flow of at least 20 sccm was maintained at all times. After setting and reaching a new pressure set point, we waited 10 min before commencing the XRD measurements to make sure that the sample fully responded to the new experimental conditions. The XRD diffractograms are background corrected by subtracting the diffractogram of an empty substrate. The values for the d111-spacing (the out-of-plane direction of the film) and the FWHM were obtained from the best fit of a pseudo-Voigt function to the background-corrected experimental data. The rocking curves were collected around the <111> diffraction peak with a Bruker D8 Discover diffractometer (Cu-K α , $\lambda = 0.1541$ nm). This diffractometer was also used for the ex situ XRD measurements. Fourier Transform Infrared Spectroscopy (FTIR): Bruker IFS66 spectrometer with KBr windows was used. The samples were mounted in a cell which can be pumped and filled with hydrogen at room temperature. A 200 nm PTFE layer was sputtered on a 10 nm Pd film on a Kbr substrate. The films was measured in the as-deposited state and after exposure to hydrogen for 10 min.

Hydrogen sensing measurements. Hydrogenation Isotherm, Kinetics and LoD Measurements: All isotherm and kinetics measurements were performed in an in-house-made vacuum chamber setup with optical windows reported earlier.^{4,21} The absolute hydrogen pressure in the chamber was monitored using two capacitive pressure gauges with different range (MKS Baratron). Optical transmittance measurements through the sample were enabled by UHV-compatible sapphire windows mounted on the vacuum chamber, and by using a fiber-coupled, unpolarized white light source (AvaLight-Hal, Avantes) and a fixed grating fiber coupled spectrophotometer (SensLine AvaSpec-2048XL, Avantes). The pressure inside the chamber was controlled using a microbar-precision leak valve. The temperature was maintained with a heating coil wrapped around the chamber and a temperature controller (Eurotherm 3216N) in a feedback loop manner, where the sample surface temperature inside the vacuum chamber was continuously used as input. All experiments were performed at constant 30 °C. The LoD measurements were performed at 1 Hz sampling frequency (0.165 ms integration time with 1000 averages). The kinetics measurements were performed at 6 Hz frequency (0.165 ms integration time with 10 averages). The observed higher noise in the kinetics data is thus a consequence of the lower number of averaged spectra per data point (Fig. S22). It is also important to note that the apparent sampling frequency obtained by the setup is slower than the theoretical one (i.e. integration time \times averaging number) due to the delay in the computing system when acquiring, processing, analyzing and plotting the data in real time. The LSPR peak descriptors were obtained by fitting a Lorentzian-function to the wavelength range at ± 60 nm around the LSPR peak in the measured optical extinction spectra, within which a good fit ($R^2 > 0.95$) is obtained, despite the asymmetry of the global LSPR peak (Fig. S19). In the experiments to determine the apparent activation energies, absorption and desorption kinetics measurements to and from 900 mbar were performed at four different temperatures, *i.e.* 30–60 °C, in 10 °C steps. For the Arrhenius analysis, t_{50} was employed as a measure of the process rate. For the long-term stability test, we stored the Pd sensors at controlled ambient conditions (average temperature 21.2 °C, humidity 23% RH, CO₂ 323 ppm). Furthermore, we intentionally performed the test on pure Pd sensors (and not on the Pd₇₀Au₃₀ alloy) for two reasons: (i) the neat Pd sensor exhibits much slower response than the analogue Pd₇₀Au₃₀ and thus its kinetics can be resolved in more detail so that any variation due to measurement artifacts can be eliminated. (ii) Neat Pd is significantly more prone to poisoning than PdAu (see Fig. S50 and also e.g. ref. 44), which makes it a more effective system to investigate the long-term stability of a polymer coating with respect to preventing sensor deactivation. Deactivation and Poisoning Tests and LoD Measurements in Ar and Air: The deactivation and LoD measurements were carried out in a quartz tube flow reactor at atmospheric pressure with optical access for transmittance measurements (X1, Insplorion AB), using synthetic air as the carrier gas, as well as Ar for the LoD experiment.⁴⁴ The gas flow rate (kept constant at 100 and 500 or 325 mL min⁻¹, for deactivation and LoD measurements in Ar and air, respectively) and gas composition were regulated by mass flow controllers (Bronkhorst ΔP). The sample inside the flow reactor was illuminated by white light (AvaLight-Hal, Avantes) through an optical fiber equipped with a collimating lens. The transmitted light was then analyzed using a fiber-coupled fixed-grating spectrometer (AvaSpec-1024, Avantes, or SensLine AvaSpec-2048XL, Avantes, for LoD measurements). The measurement temperature was maintained at 30 °C. Quartz Crystal Microbalance Measurements: The QCM window-module (QSense Explorer Microscopy, Biolin Scientific Corporation) was connected to a series of mass flow controllers (Bronkhorst ΔP) to regulate the H₂ partial pressure in Ar carrier gas at a constant total flow rate of 30 mL min⁻¹. Data collection was executed by QSoft® (Biolin Scientific AB). The absorbed hydrogen content in the nanoparticles was calculated from the QCM frequency shift via the Sauerbrey equation⁴⁵ and analysis of the nanoparticle surface coverage based on SEM images of the crystal, as described in detail elsewhere.²² All of the measurements were carried out at 30 °C.

Density Functional Theory calculations. All the modeled structures were optimized within the DFT framework. The cluster models were optimized using the PBE functional⁴⁶ while van der Waals contributions were included for the polymer and polymer-H₂ systems by using the vdW-CX functional.⁴⁷ Geometry optimizations were performed using the Vienna ab initio simulation package, VASP.^{48,49} Projector augmented waves (PAW)^{50,51} were used, with a plane-wave kinetic energy cutoff of 500 eV. To avoid artificial interaction due to the periodic boundaries conditions, a $30 \times 30 \times 31$ Å supercell was used, ensuring at least 16 Å of vacuum between two successive images. Γ -centered⁵² calculations were performed with a Gaussian smearing, σ , of 0.02 eV, with the energies being extrapolated to $\sigma = 0.00$ eV. The atoms positions were optimized until the criterion of the residual forces on any direction being less than 0.02 eV Å⁻¹ was met.

Finite-Difference Time-Domain simulations. We used FDTD Solutions to calculate the optical spectra of a Pd nanodisk as a function of the refractive index of the coating, as well as coating thickness. Due to the fact that the fabricated arrays are amorphous in terms of the nanoparticle arrangement on the surface, we cannot model exactly the whole sensor and accordingly are limited to either a single disk or, to account for inter-particle coupling, a periodic lattice. In the

present case we first chose the former solution, although in one case we also consider a hexagonal lattice. The Pd nanodisk is modeled with a diameter of 180 nm, 30 nm thickness and rounded edges. The permittivity of Pd and Pd hydride (PdH_{0.67}) are taken from the literature.⁵³ The structures are placed directly on top of a glass substrate (n = 1.45) and are conformally coated by an isotropic dielectric layer with a variable thickness and refractive index (RI). In the simulations, we span the thickness in the range from 5 to 75 nm and the RI from 1 to 1.6. This encompasses the range of the parameters considered in the experiment. The structures are illuminated by a plane wave and extinction is collected and analyzed in terms of resonance position.

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Author contributions

F.A.A.N. and C.L. designed the experiments, analyzed the data and wrote the manuscript. F.A.A.N. and I.D. fabricated the sensors. F.A.A.N. performed sensing measurements on PTFE sensors. F.A.A.N. and I.D. performed sensing measurements on PMMA and tandem sensors. L.C. and A.H. executed the DTF calculations. A.S.-A. performed the XPS analysis. H.S. deposited the PTFE thin films. L.J.B. and B.D. performed the XRD analysis. A.B.d.S.F. and J.B.W. performed the TKD analysis. S.K. performed the STEM-EDS analysis. T.J.A. performed the FDTD simulations. V.P.Z. contributed the theoretical analysis on the sensor kinetics and PTFE strain. B.D. and C.L. coined the initial idea. C.L. coordinated the project.

Additional information

Supplementary Information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints.

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Competing financial interests

The authors declare no competing financial interests.