Computational Design of Alloy Nanostructures for Optical Sensing of Hydrogen

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Cite This: https:/	/doi.org/10.1021/acsanm.2c01	189	Read Online	
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ABSTRACT: Pd nanoalloys show great potential as hysteresis-free, reliable hydrogen sensors. Here, a multiscale modeling approach is employed to determine optimal conditions for optical hydrogen sensing using the Pd–Au–H system. Changes in hydrogen pressure translate to changes in hydrogen content and eventually the optical spectrum. At the single particle level, the shift of the plasmon peak position with hydrogen concentration (i.e., the "optical" sensitivity) is approximately constant at 180 nm/ $c_{\rm H}$ for nanodisk diameters of $\gtrsim 100$ nm. For smaller particles, the optical sensitivity is negative and increases with decreasing diameter, due to the emergence of a second peak originating from coupling between a localized surface plasmon and interband transitions. In addition to tracking peak position, the onset of extinction as well as extinction at fixed wavelengths is



considered. We carefully compare the simulation results with experimental data and assess the potential sources for discrepancies. Invariably, the results suggest that there is an upper bound for the optical sensitivity that cannot be overcome by engineering composition and/or geometry. While the alloy composition has a limited impact on optical sensitivity, it can strongly affect H uptake and consequently the "thermodynamic" sensitivity and the detection limit. Here, it is shown how the latter can be improved by compositional engineering and even substantially enhanced via the formation of an ordered phase that can be synthesized at higher hydrogen partial pressures.

KEYWORDS: hydrogen sensing, nanoplasmonics, localized surface plasmon resonance, nanoparticles, palladium alloys, dielectric function

INTRODUCTION

One of the main challenges of the hydrogen economy, a future energy system where fossil fuels have been replaced with hydrogen-based fuels, is the flammability of hydrogen gas under ambient conditions.¹ As a result, hydrogen sensing continues to be a very active research field with the goal of providing fast, reliable, and long-term stable hydrogen sensors that can prevent major accidents. Several different sensing platforms have been proposed, typically based on the change in optical²⁻¹² or electrical¹³⁻¹⁵ properties of a material during hydrogen absorption. Many of these devices are based on palladium (Pd), which forms a hydride phase upon exposure to a hydrogen-rich atmosphere. The electronic properties of the hydrogenated system differ from the hydrogen-free one, leading to a shift in the optical response as well as a change in resistivity, which can be measured and related to the hydrogen pressure in the surrounding environment. In this work, we focus on optical sensors based on the localized surface plasmon resonance (LSPR) of Pd-Au-based nanoalloys^{2,3,6,9} and show how they can be optimized via geometry and alloy composition.

A significant shortcoming of hydrogen sensors based on pure Pd nanoparticles (NPs) is that they undergo a first-order phase transition upon hydrogenation, from the hydrogen-poor α to the hydrogen-rich β -phase, which leads to a nonlinear, discontinuous response to hydrogen pressure. The phase transition is further associated with significant hysteresis,¹⁶ which ultimately makes the response of the sensor dependent on its history. It has been shown that these disadvantages can be overcome by alloying.^{17,18} Specifically, the introduction of about 20% of gold (Au) suppresses the phase transition, making the response of the sensor a linear function of hydrogen pressure.^{6,17,19-21} To optimize these systems for hydrogen sensing, important questions remain: in what proportions should we mix the two alloyants, what shapes should the NPs have, and what features of the optical response should the sensing mechanism be based on? Historically, similar questions have been addressed by trial-and-error experiments. While that is a viable approach, the additional

Received: March 17, 2022 Accepted: July 8, 2022



Scheme 1. Methodological Overview^a



degrees of freedom introduced by mixing two or more components and the large space of available geometries make it extraordinarily cumbersome to experimentally test all combinations of potential interest. In this context, a multiscale modeling approach that resolves the variation in electronic properties with composition and translates this variation to the optical properties of NPs of different sizes and shapes is therefore invaluable.

The dielectric function (DF) links electronic and optical properties, as well as atomic and continuum scales. We have recently calculated the DFs of 10 binary alloys in their full composition range and demonstrated that electrodynamic simulations based on these DFs provide accurate predictions of the optical response of alloy nanoparticles.²² Earlier studies of pure Pd have shown, in principle, how one can combine electronic structure calculations and electrodynamic simulations to enable multiscale modeling of the optical response during hydrogenation.^{23,24} They were, however, limited in the representation of chemical order (due to very small structure models) and also did not consider Pd-alloys. Here, we build on and extend the underlying approach to calculate the DFs of Pd-Au alloys with up to 46% Au and in the full range from no hydrogen to full hydrogen loading. The DFs are then converted to a Lorentzian representation and used in electrodynamic simulations to predict the extinction spectra of Pd–Au nanodisks (Scheme 1). Finally, the optical sensitivity of hydrogen sensors based on Pd-Au NPs is derived from the changes in their extinction spectra. To obtain the actual sensitivity one must also include a thermodynamic factor that accounts for the solubility of hydrogen in the material, which here is taken into account using a thermodynamic model developed previously.²⁵ This multiscale approach, which extends from first-principles calculations on the atomic scale to electrodynamic simulations on the macroscale, is not limited to hydrogenated Pd-Au nanoparticles but can in principle be applied to the optical response of any material.

An important aspect in this context is atomic scale ordering, which has the potential to significantly alter the DF and, in turn, the optical response.²² This situation requires careful consideration when calculating DFs from first-principles and applying them to the prediction of the optical response of nanoalloys. In thermodynamic equilibrium and in the absence of hydrogen, bulk Pd–Au is expected to form an alloy without long-range order and at most a low degree of short-range

order.^{25,26} Furthermore, since in many studies the nanoalloys are annealed at high temperatures, 6,17,21 it is safe to assume that the chemical distribution of the alloyants is close to random. To represent this situation in our calculations, we use special quasi-random structures (SQSs),²⁷ which are constructed to reproduce the (lack of) short-range order in a truly random alloy using unit cell sizes that are small enough for efficient calculations (in our case 24 Au/Pd atoms and 0-24 H atoms). As discussed before,^{25,26,28} exposure to hydrogen can, however, impact thermodynamics and ordering. As the nanoalloys are typically only exposed to hydrogen at high temperatures during fabrication and for short periods of time during operation, it is reasonable to assume that this effect is negligible for the samples produced in, for example, refs 6, 17, and 21. It is, however, known from both experimental and theoretical evidence that an ordered, intermetallic phase can emerge for compositions around 25% Au after annealing at high hydrogen pressures and moderate temperatures.^{25,26} As we will show in this work, the presence of this phase could potentially improve the sensing ability at low hydrogen pressures radically.

RESULTS AND DISCUSSION

Electronic Structure and Dielectric Functions. We begin by examining the DFs as calculated with linear-response time-dependent density-functional theory (Figure 1a,b; see Figure S1 and Figure S2 for the real and imaginary parts at more compositions and in a wider energy range). A few trends are apparent.

In the case of Pd–H, addition of H leads to the emergence of two peaks in the imaginary part of the DF that at 100% H loading are located around 1.5 eV and at approximately 3.0– 3.5 eV (Figure 1a). Comparison with the density of states (DOS) indicates that these two features are related to two pronounced peaks in the d band (Figure 1c; see Figure S3 for the DOS at more compositions). In fact in this case, the DOS in its entirety is almost unaffected by H addition, save for a rigid shift of the energy scale. This is a result of the localized character of the H states, which do not hybridize with each other. Addition of H also reduces the magnitude of the real part of the DF in the visible range, as has been previously observed in both experiment^{29,30} and computations²⁴ (Figure S4). Generally speaking, such a trend is expected to imply a

Figure 1. Variation of the imaginary part of the dielectric function as a function of photon energy for (a) PdH_x without Au and (b) $Pd_{1-y}Au_y$ in the absence of H, as calculated in this work. The dielectric function is closely related to the density of states (c, d). Energies have been shifted such that the Fermi level is always at 0 eV.

redshift of the plasmon peak, as the Fröhlich condition for LSPR emergence is satisfied at lower energies.

Upon addition of Au to Pd, there is almost no change in the imaginary part of the DF in the composition range considered here (Figure 1b). In the DOS, on the other hand, a notable smoothing occurs as a result of hybridization between the d states of Pd and Au (Figure 1d).³¹ Therefore, when H is added to Pd-Au, the two peaks that were apparent in the DF of Pd-H are largely smoothed out due to alloy disorder (Figure S2).

Optical Response. After fitting the DFs to a Lorentzian representation (as described in Supplementary Note 1) we carry out electrodynamic simulations of the optical response of truncated cone nanodisks (Figure 2a), characterized by their height h (20–40 nm) and aspect ratio (AR) (2–12) or diameter $d = AR \times h$ on a silica substrate, mimicking a single nanodisk fabricated by the hole-mask colloidal lithography method.³² The exact shape of the nanodisk is, however, not of great importance (see Figure S5 for a comparison between slightly differently shaped nanodisks).

Pd–H Nanodisks. We begin by studying the optical response of the Pd–H system based on the calculated extinction. In Figure 2b–e the extinction spectra for four specific geometries are shown (see Figure S6 for separate scattering and absorption spectra). These are selected since they represent different characteristics of the studied spectra (see Figure S7 for the full range of geometries). The shape of the extinction spectra varies with geometry and H content, but at least one peak can be identified for all systems. Most geometries considered have extinction spectra similar to Figure 2b with a well-defined, relatively sharp peak that shifts to the

Figure 2. Optical response of truncated cone nanodisks of varying height *h* and diameter *d* and fixed cone angle $\theta = 60^{\circ}$ (a). Here, we show the extinction spectra for disk geometries $(d \times h \text{ in nm})$: 240 × 30 (b, f), 90 × 30 (c, g), 80 × 40 (d, h), and 40 × 20 (e, i) with 0% Au (b–e) and 25% Au (f–i) at H concentrations 0–67%. The dashed lines correspond to H concentrations $c_{\rm H} > 0.67 - c_{Au}$, which are excluded in the calculations of the optical sensitivity \tilde{S} .

red with increasing H content. For smaller nanodisks, however, the features are broader and we can identify two distinct peaks at H concentrations above 50% (Figure 2c-e). The second peak emerges with increasing H content and decreasing height and diameter and becomes the dominant feature in the high H loading limit for the smallest particles considered here.

The high-energy feature is in fact present for all geometries in the high H limit, but for larger nanostructures (such as the one in Figure 2b) this feature is less noticeable due to the larger amplitude of the low-energy peak. Of these two peaks, the high-energy one is roughly constant in energy (and amplitude; see Figure S8) while the low-energy peak shifts when the aspect ratio is varied. To illustrate this behavior, we compare the cases of 0% and 100% H in Pd, which allows one to identify the key features in the DFs more easily due to the lack of disorder on the H sublattice. For 0% H in Pd, we observe one peak that shifts from 1 eV for large nanodisks to 5 eV for small nanodisks (Figure 3a), in agreement with the expected size dependence of a LSPR. For 100% H in Pd, on

Figure 3. Peak position of the two peaks in the extinction spectra for PdH (opaque lines) and Pd (transparent lines) as a function of aspect ratio for two disk heights h (a) and the corresponding dielectric functions (b). PdH displays two features, one low-energy peak (solid line) which shifts with aspect ratio and one high-energy peak (dashed lines) which only shifts for small aspect ratios, indicating avoided-crossing behavior due to coupling between the peaks. The high-energy peak can be traced to the feature at 3.5 eV in the dielectric function (b) for 100% H (opaque line). This feature is not present for 0% H (transparent line), and as a result, the LSPR peak can shift over a much wider energy interval (transparent lines in b). Note that (b) shows the fitted Lorentzian representation of the dielectric functions.

the other hand, we observe two peaks (see the corresponding extinction spectra and field enhancement in Figure S9). For nanodisks with larger ARs, the peaks are separated in energy and the low-energy peak increases in energy with decreasing AR (as expected for a LSPR) while the high-energy peak remains at about 3.5 eV (Figure 3a). The lack of size dependence for the high-energy peak indicates that it is caused by a bulk phenomenon, such as a spectrally localized interband transition.^{33,34} When a LSPR peak moves closer to an interband transition, they can couple and exhibit avoided crossing.^{33,34} This is precisely the behavior we observe for small ARs when the shift of the low-energy feature is restricted (compared to the 0% H case) and the high-energy peak starts to increase in energy. The presumed interband transition can be related to the previously discussed feature in the DF close to 3.5 eV (Figure 3b and section Electronic Structure and Dielectric Functions). For 0% H in Pd there is no such feature, and as a result, there is no avoided crossing for 0% H and the LSPR peak can shift over the entire energy interval without interference.

Pd-Au-H Nanodisks. We now turn to the impact of Au on the extinction spectra. For comparison with previous experimental studies, 6,21 we focus on 25% Au which provides a trade-off between suppressing the hysteresis and avoiding the decrease in H absorption with increased Au concentration (Figure S10). The introduction of 25% Au has a relatively small effect on the optical response (Figure 2). Some changes in the extinction spectra can, however, be identified. For the most part, introducing Au leads to a broader extinction spectrum with slightly lower peak amplitude and a weaker tendency for peak splitting (Figure 2g-i). This is consistent with the observation that the DOSs and the DFs are typically smoothed by adding Au to Pd; features are blurred out by the chemical disorder. In the case of large AR and high H content, however, the LSPR peak becomes sharper and more intense with increasing Au content (Figure 2f). This sharpening is a consequence of the increased distance between d band and Fermi level in systems with high Au and/or H content (Figure 1c), which reduces the ability of the d electrons to screen the LSPR at low energies. Peak splitting occurs in the high H limit for Pd-Au nanodisks as well, but the evolution of the high-

Figure 4. For a given set of extinction spectra, such as the spectra (a) without or (b) with pronounced peak splitting, one can define the optical sensitivity \tilde{S} based on several different shifts with H content (illustrated here for Pd–H). First, the peak position (black dots in (a) and (b)) generally shifts linearly with H concentration (c). For the case of pronounced double peaks, we study the peak shift both excluding data points with double peaks (solid lines in (c)) and with double peaks (dashed lines in (c)). In the latter case, only the data points corresponding to the two lowest and two highest H concentrations (indicated by the shaded regions) contribute to the fit. Second, the wavelength at a certain onset amplitude (50% of the peak extinction at 0% H here) has a similar linear increase with increased H content (d). Third, the extinction at a particular wavelength (900 nm here) also shows an almost linear dependence on the H concentration (e), although the trend deviates slightly from linearity close to the peak(s). In all these cases, the optical sensitivity \tilde{S} can be defined as the slope of the linear trend(s).

illustration, can be written as

energy peak with size or H content is considerably less regular compared to the case of Pd. This is related to the corresponding somewhat irregular evolution of the alloy DFs which is caused by the imperfect representation of the chemical order due to limited system size.

Optical Sensitivity Optimization. In general, sensitivity is defined as the ratio between the output signal and the underlying external stimulus. In *optical* sensing it is common to follow the variation of one or more characteristic descriptors of the optical spectrum as measures of the output signal, which in the present case of plasmonic sensing are related to the behavior of the plasmon peak, e.g., its amplitude, width or position, or the surrounding spectral region. In the specific case of hydrogen sensing the external stimulus is the presence of H₂ in the surrounding environment as expressed by its partial pressure $p_{\rm H_2}$. More formally the sensitivity of such a sensor, using here the position of the plasmon peak $\lambda_{\rm max}$ for

$$S = \frac{d\lambda_{\max}}{dp_{H_2}} \tag{1}$$

Earlier measurements for this material system^{2,3,6,9} have, however, commonly considered the change in the absorption maximum λ_{max} with the hydrogen concentration in the material c_{H} ,

$$\tilde{S} = \frac{d\lambda_{\max}}{dc_{\rm H}} \tag{2}$$

The two measures are related to each other via

$$S = \frac{d\lambda_{\max}}{dc_{\rm H}} \times \frac{dc_{\rm H}}{dp_{\rm H_2}} = \tilde{S}S_{\rm sol}$$
(3)

where the subscript emphasizes that S_{sol} directly depends on the solubility of hydrogen as a function of its partial pressure.

In the present section, we will focus entirely on \tilde{S} , for brevity referred to as the optical sensitivity. We will return to the role of S_{sol} in the section Thermodynamic Optimization.

In the following, we study three specific measures for the optical sensitivity \tilde{S} based on the shift in peak position (*peak shift*), wavelength at a fixed extinction amplitude onset (*onset shift*), and extinction at a certain wavelength (*extinction shift*), respectively (Figure 4), with respect to the absorbed H concentration to determine the limits of optical sensitivity optimization through engineering of composition and/or geometry.

In the case of Pd–H, the H concentration range that is relevant under common thermodynamic conditions is rather wide, while it is more narrow for Pd–Au–H. As a consequence, we have chosen to restrict the maximum H concentration considered to $c_{\rm H} < 0.67 - c_{\rm Au}$ when calculating the sensitivities, which is sufficient for hydrogen pressures up to 1 bar (Figure S10). The exact limit is not of great importance since the studied shifts are generally linear with H concentration for hydrogen pressures of ≤ 1 bar.

Peak Shift. In ref 6, the (optical) sensitivity was defined as the ratio of the peak shift to the change in H concentration. For a specific geometry, the peak position is generally an approximately linear function of the H concentration (Figure 4c), and the sensitivity can be obtained as the slope of the corresponding linear fit. This is straightforward for the larger

nanostructures considered (i.e., disks well-represented by Figure 4a) that feature one dominant peak for all H concentrations.³⁵ For the smaller nanodisks considered here, peak splitting, however, comes into play which complicates the determination of the optical sensitivity. In systems with significant peak splitting (e.g., see Figure 4b), we therefore consider two different peak shift sensitivities. First, we fit the peak shift as a function of H concentration after excluding all H concentrations with peak splitting (solid line in Figure 4c). Note that with this approach, cases with pronounced peak splitting typically result in a large fitting error since the data points are characterized by large scatter (see Figure S13 for the fitting errors). Second, we fit the peak shift based on only the first two data points (low H limit) and the last two data points (high H limit) for either peak separately (dashed lines in Figure 4c), resulting in two sets of sensitivities, typically with opposite signs. The latter approach is suitable for pure Pd and low Au content since the phase transition introduces a jump from low to high hydrogen content (Figure S10). For high Au content ($\geq 10\%$), peak splitting is in practice not an issue since the hydrogen content is below 50% at relevant pressures (Figure S10).

In Figure 5a, we display the obtained peak shift sensitivity for all considered systems as a function of the disk diameter. Clearly, the size dependence of the sensitivity can be divided into two regimes. Nanodisks with diameter of ≥ 100 nm have a roughly constant sensitivity around 180 nm/(H/(Au + Pd)), which to a large degree is independent of disk height and alloy composition. Smaller nanodisks, however, follow a linearly decreasing trend with increasing diameter. The double peak sensitivities (semicircles in Figure 5a) follow the same trends and introduce an overlap between the two regimes. The trend for small nanodisks can be explained by the peak splitting. When the high-energy peak at 67% H ($\lambda_{H_{max}}$) dominates over the corresponding low-energy peak, the sensitivity is determined by the shift between the 0% H peak (λ_0) and the former. In a simplified picture, λ_0 shifts to lower wavelengths with decreasing diameter while $\lambda_{H_{max}}$ is constant. This means that, as the diameter is decreased, the sensitivity will shift from strongly negative to positive, crossing zero when λ_0 and $\lambda_{H_{max}}$ align.

The optical sensitivity trends are largely unaffected by the introduction of up to 25% Au (Figure 5a). This is somewhat surprising since for Pd the sensitivity is almost entirely determined by peak splitting and for Pd—Au the peak splitting is less distinct and often not relevant due to the H concentration threshold. Clearly, the peak shift sensitivity is not very sensitive to the details in the shape of the extinction spectra.

In Figure 5a, we include experimental results from ref 6, where the authors find a linear increase in the (optical) sensitivity as a function of the peak position at 0% H.³⁶ While the experimental trend can be interpreted to reach a plateau for peak positions at 0% H of $\gtrsim 600$ nm, there is still an apparently large discrepancy with our findings. Note, however, that the experimental results were obtained for nanodisk *arrays* and the measurements thus correspond to an average over a large *ensemble* of particles. Notably, the measured (optical) sensitivity is very close to zero right at the discontinuity between the two sensitivity regimes, where peak-splitting is most pronounced. This could suggest that the experimentally measured linear increase in sensitivity is due to a superposition

Figure 5. Optical sensitivity of Pd–Au nanodisks with varying Au content as a function of the disk diameter, in terms of (a) peak shift, (b) onset shift, and (c) extinction shift at λ_{opt} per unit H uptake. In (a), semicircles correspond to the sensitivity when peak-splitting is taken into account (as explained in Figure 4) and plus signs show experimental data from ref 6. Note that the experimental sensitivity is rescaled from nm/(H/Pd) to nm/(H/(Au + Pd)) and the disk diameter is approximated by the linear relation in Figure S12b. The corresponding peak position at 0% H is indicated by the upper *x*-axis scale, calculated from the diameter and the linear relation in Figure S12b. In (c), the inset shows the wavelength where the optimal sensitivity was found for each system.

of nanoparticle shapes and sizes that fall on either side of the discontinuity between the sensitivity regimes, and as the fraction of particles which display peak splitting decreases, the sensitivity increase. We address this aspect in detail in the section Uncertainty Analysis but find that size distribution and array effects on their own are insufficient to explain this effect.

Onset Shift. Another way of defining the optical sensitivity, which circumvents keeping track of peaks, is to monitor the wavelength at the onset of a fixed target extinction amplitude. Here, we define the target amplitude as 50% of the peak amplitude at 0% H (Figure 4a,b) and track the corresponding wavelength (to the right of the peak in wavelength) as a function of the H concentration (Figure 4d).³⁷

The onset shift generally follows a linear trend with respect to H concentration, the slope of which can be defined as the sensitivity (Figure 5b). For almost all considered geometries, such as the ones in Figure 4a,b, we obtain a positive sensitivity that increases slightly with increasing diameter. For very small nanostructures, however, the low energy peak at high H content falls below the onset amplitude (Figure 2e,i), which reverses the direction of the onset shift and results in a small or negative sensitivity. In principle, this issue could be resolved by decreasing the onset amplitude value below 50%. That would, however, push the amplitude onset for the largest nanodisks beyond the low energy (high wavelength) limit.

The introduction of Au slightly reduces the slope of the sensitivity trend such that the sensitivity of the largest nanodisks decreases with increased Au content.

Extinction Shift at a Certain Wavelength. A third possible optical sensitivity definition is based on the shift in extinction amplitude at a certain wavelength (Figure 4a,b,e). This definition is of particular interest for real devices since it does not require measuring the spectrum over the entire spectral range but only at a single wavelength. In Figure S14, we display extinction spectra at increasing H content in terms of extinction at selected energies (Figure S14a-d) and the corresponding slope of linear fits (Figure S14e-h).

The slope changes significantly with energy from a positive value to the left of the peak to a negative value to the right of the peak. This means that the choice of wavelength is crucial for maximizing the optical signal. For each system we define the wavelength for which the largest slope (in absolute value) is obtained as λ_{opt} . The extinction shift sensitivity is defined as the absolute value of the slope at λ_{opt} in units corresponding to extinction cross section (Figure 5c). The sensitivity increases with disk diameter, decreases slightly with Au content, and in the size range considered is independent of disk height. It should be noted that, in contrast to peak and onset shift, the extinction shift is proportional to the amplitude of the optical response of an individual nanodisk. As a result, the sensitivity increase is to a large degree caused by the increase in extinction cross section with size (Figure 2). In a real device, the number of nanodisks per area can be increased for smaller particles which means that for some purposes, it might be more relevant to compare the extinction-shift sensitivity scaled with nanodisk area.

Uncertainty Analysis. As previously noted, there is a discrepancy between our results and an earlier experimental study^{\circ} for the peak shift sensitivity. Such differences between experiment and simulations are not unexpected given the many uncertainties involved, which exist both at the level of the electrodynamic simulations of the optical response and at the level of the electronic structure calculations of the DFs. Here, we investigate the impact of the sources of discrepancy that we judge to be the most important.

Let us first consider the effect of particle size distribution and array effects on the optical spectrum, which enter at the level of the electrodynamic simulations. In contrast to the single particles simulated in this work, the experimental samples consist of many irregularly shaped nanoparticles with a significant size distribution.⁶ These particles are furthermore randomly distributed on a substrate, leading to array effects that are known to influence the optical spectrum.³⁹ We observe that the naive approach of weighting extinction spectra according to some size distribution does not significantly alter the extinction (Figure S15). To better mimic the experimental situation, we therefore performed additional simulations of random arrays of nanoparticles (using the T- matrix method as described in the section T-Matrix Simulations). While the array effects (but to a lesser extent the size distribution) do change the position and sharpness of the peak (Figure S16; also see ref 40), the sensitivity trends remain similar to the single particle results (Figure 6).

Figure 6. Optical peak shift sensitivity \hat{S} for Pd nanodisks with 30 nm diameter calculated using different computational setups and DFs. The data from Figure 5a are included in gray. The sensitivities for explicit and effective random arrays (orange), with and without size distribution, follow closely the previously discussed trends for single particles (large gray (semi)circles). The sensitivities obtained using DFs from Silkin et al.,²⁴ Palm et al.,²⁹ and Vargas et al.³⁰ show a large spread. Note that, except for the data in gray, the sensitivities here are calculated based on only the peaks at 0% and ~67% H ³⁸ in order to reduce the computational effort.

A further source of uncertainty is the DFs, which in the present work are obtained from electronic structure calculations. The calculated DF depends on the underlying electronic structure method, in the present case primarily the exchange-correlation functional, but also on the treatment of scattering, especially due to defects (e.g., including surfaces, grain boundaries, and dislocations) and chemical disorder in the case of alloys. With regard to the description of the electronic band structure, the present approach yields results that are in very good agreement with recent highly accurate many-body theory calculations.⁴¹ We therefore focus our attention on the possible effect of scattering mechanisms. In Figure 6, we compare the optical sensitivity calculated using three sets of DFs from the literature; two experimentally measured ones (Vargas et al.³⁰ and Palm et al.²⁹) and one calculated using an approach similar to ours (Silkin et al.²⁴). Although the DFs are in semiquantitative agreement over the energy range considered (Figure S4), small differences clearly can have a large effect on the resulting sensitivity. Note in particular the large span of sensitivities resulting from the experimentally measured sets of DFs, while the calculated set of DFs yield sensitivities very similar to our original findings. Real samples are typically subject to defect scattering which leads to a broadening of the Drude peak. This effect is very difficult to include rigorously at the ab initio level and is furthermore sample-dependent, which could explain the large difference between the two experimentally measured sets of DFs. To

illustrate this aspect further, it is instructive to consider the case of the noble metals, for which the d-band feature in the DFs is clearly separated from the Drude peak and thus allows one to observe the effect of defect scattering more directly. This reveals the large variation in the width of the Drude peak caused by differences in sample preparation and the resulting defect density (e.g., see Figure S1a-c of ref 22). In the case of Pd alloys and hydrides the d-band overlaps with the Fermi level, which leads to a very broad response in the low-energy region and the impact of (defect) scattering cannot be clearly separated. This makes it difficult to include this contribution systematically at the level of the DFs and eventually assess its impact on the optical response.

Regardless of which set of DFs is used, a large shift in sensitivity appears when the peak position at 0% H is at approximately 450 nm, which is also where the experimentally measured sensitivity goes to zero, in line with the discontinuity between the two sensitivity regimes previously discussed. This indicates that it is in fact the interband transition at high H content that is responsible for large changes of the sensitivity. We verified this argument by explicitly removing the associated Lorentzian from the DF, which results in a sensitivity that remains at a high, positive value even as the peak position at 0% moves below 450 nm (Figure S17).

Lastly, we observe that by introducing artificial Lorentzian broadening in the simulated extinction spectra, a linear increase in the sensitivity similar to the experimental results can be obtained (Figure S18), although quite extreme levels of broadening are necessary. We conclude that it is in general very difficult to account quantitatively for the sample-specific broadening channels.

Thermodynamic Optimization. Above, in agreement with earlier experimental studies, we have seen that the alloy composition is not crucial for the *optical* sensitivity \tilde{S} . It is, however, crucial for the *thermodynamic* sensitivity S_{sol} and hence impacts the actual sensitivity S see eq 3. In this section, we therefore analyze the dependence of the *thermodynamic* sensitivity S_{sol} on alloy composition and then translate the results to a detection limit in terms of the hydrogen pressure. In this context, we also show that the highest sensitivities are obtained for the ordered L1₂-Pd₃Au phase.

Thermodynamic Sensitivity. For the following analysis, we limit us to a temperature of 300 K and use the relationships between partial pressure of H_2 , Au concentration, and H concentration established in ref 25 by means of alloy cluster expansions, Monte Carlo simulations, and experimental data (Figure 7a and Figure S10). Using these data, one can compute the (differential) thermodynamic sensitivity (also see eq 3),

$$S_{\rm sol} = \frac{dc_{\rm H}}{dp_{\rm H_2}} \tag{4}$$

In practice one is, however, more often interested in the change relative to a reference pressure, representing a situation in which one aims to probe a sudden increase in hydrogen pressure relative to a low background level. Here, we therefore consider the thermodynamic sensitivity defined as

$$\tilde{S}_{\rm sol} = \frac{c_{\rm H}(p_{\rm H_2}) - c_{\rm H}(p_{\rm H_2}^{\rm ref})}{p_{\rm H_2} - p_{\rm H_2}^{\rm ref}}$$
(5)

Figure 7. (a) Hydrogen partial pressure as a function of hydrogen concentration at a temperature of 300 K for different gold concentrations according to ref 25. The dashed lines indicate the miscibility gap between the H-poor α -phase and the H-rich β -phase. (b) Thermodynamic sensitivity \tilde{S}_{sol} according to eq 5 using a reference pressure of $p_{H_2}^{ref} = 10^{-9}$ bar as a function of Au concentration at different H₂ partial pressures and a temperature of 300 K.

While there are some quantitative differences between these two measures, they show qualitatively the same behavior (Figure 7 and Figure S11).

The thermodynamic sensitivity varies with both Au concentration and H₂ partial pressure (Figure 7b). While at smaller partial pressures (≤ 1 mbar) one obtains a monotonic *increase* of the thermodynamic sensitivity with Au content, at larger partial pressures (≥ 20 mbar) one actually observes a *decrease* of \tilde{S}_{sol} with Au content. This behavior can be readily rationalized in terms of the pressure–concentration isotherms (Figure 7a). In the intermediate region around approximately 10 mbar, one observes a maximum, which shifts with hydrogen partial pressure.

Since the actual sensitivity *S* is simply the product of the optical sensitivity \tilde{S} and the thermodynamic sensitivity S_{sob} see eq 3, and the optical sensitivity does not strongly depend on Au concentration, the behavior described above should dominate the composition dependence of the actual sensitivity. One should, however, also take into account the fact that experiments are commonly carried out for ensembles of particles, for which the p_{H_2} - c_{H} isotherms are considerably broadened. As a result, the variation of both the thermody-

namic sensitivity $(S_{sol} \text{ or } \tilde{S}_{sol})$ and the actual sensitivity (S) with Au concentration should be less pronounced than in the present case, which models the thermodynamic behavior of single particles under idealized conditions.

Detection Limit. Next, we translate our results to a detection limit in terms of hydrogen pressure. We assume a fixed optical sensitivity \tilde{S} as defined by the gray, dashed line in Figure 5a (180 nm/(H/(Pd + Au))). We then explore the hydrogen pressure required to induce a certain LSPR peak shift (Figure 8a). In stable, experimental setups, a peak shift of 0.1

Figure 8. (a) Hydrogen pressure required to induce a peak shift of 10 nm (blue), 1 nm (orange), and 0.1 nm (green) at 300 K assuming a randomly ordered Pd–Au sublattice (dots) or $L1_2$ ordering (stars). Peaks shifts are calculated from the dashed, gray line in Figure 5a (180 nm/(H/(Au + Pd))) based on the relation between H₂ partial pressure and Au/H concentration calculated in ref 25. The flammability limit and the lower target for the detection limit of hydrogen sensors set by the US DOE⁴² are marked in dashed red and gray, respectively. (b) Temperature dependence for the detection limit assuming a peak shift of 1 nm can be resolved for random Pd_{0.75}Au_{0.25} (orange) and the same composition but with L1₂ ordering (black).

nm or less can be detected fairly easily. This implies that the detection limit is well below 1 mbar (green line in Figure 8a), which is approximately the target (1000 ppm) set by the U.S. Department of Energy (DOE) for hydrogen sensors⁴² (dashed, gray line in Figure 8a). This is true regardless of Au content in the concentration interval studied here. Under more challenging circumstances, however, it cannot be ruled out that a more significant peak shift is required to obtain a reliable signal. With a 1 nm peak shift (orange line), nanodisks of pure

Pd do not meet the U.S. DOE target, and at 10 nm (blue line), the target cannot be met at all.

Enhancing Sensitivity through Chemical Ordering. The situation can be substantially improved if we consider the ordered $L1_2$ -Pd₃Au phase, which is expected to form when the alloy is subjected to high pressures of H_2 .²⁵ This compound absorbs large amounts of hydrogen already at pressures below 10^{-5} bar, for which the regular alloy shows almost no sorption (Figure 7a), as observed in both experiments²⁶ and calculations²⁵ (also see Figure S10). As a result, one obtains a substantially larger thermodynamic sensitivity than for the regular alloy, especially for pressures of $\lesssim 1$ mbar (stars in Figure 7b).

To evaluate the suitability of this $L1_2$ -Pd₃Au compound for optical hydrogen detection, we calculated the dielectric function at different hydrogen contents and performed electrodynamic simulations in the same fashion as for the disordered phase (Figure S19). The results are very similar for the ordered and the disordered phase, with almost identically positioned LSPR peaks. We can thus assume the same sensitivity with respect to the peak position as for the disordered phase. The larger variation of the hydrogen uptake with pressure translates, however, to a detection limit that is predicted to be approximately 1–2 orders of magnitude lower in the ordered compound compared to the disordered phase, regardless of whether a large or small peak shift can be detected (stars in Figure 8a).

When temperature is increased, a higher hydrogen pressure is required to maintain the same hydrogen content in Pd–Au. Consequently, a higher hydrogen pressure is required to achieve a certain peak shift. The detection limit thus decreases with increasing temperature. The hydrogen uptake is, however, significantly higher in the L1₂-ordered phase than in the random phase, also at higher temperatures (Figure 8b). Hydrogen sensors based on the L1₂-ordered phase may thus retain a satisfactory detection limit at temperatures at which random alloys no longer meet the requirements.

CONCLUSIONS

Our results indicate that for all optical sensitivity measures, the disk diameter is the most relevant descriptor (Figure 5) in the selected span of geometries while disk height, AR, and alloy composition are of secondary importance. The highest optical sensitivities are found for larger particles, but for disk diameters above 150 nm the improvement is limited. For the optical sensitivity based on peak shift, our results reveal two separate regimes with respect to particle size (Figure 5a). For larger nanodisks ($d \ge 100$ nm), the optical sensitivity is approximately constant at 180 nm/ $c_{\rm H}$. For smaller nanodisks ($d \leq 100$ nm), on the other hand, the optical sensitivity starts at a negative value and increases with decreasing particle size. This behavior originates from the emergence of a second peak that appears for H concentrations above 50%, caused by a localized interband transition, which becomes more apparent as the particle dimensions shrink.

The peak shift sensitivities obtained from electrodynamic simulations using the calculated DFs are comparable in magnitude to direct experimental measurements. They are, however, consistently larger and more importantly in the region where direct comparison is possible remain constant, whereas the experimental data show a linear shift with the absorption maximum. According to our calculations, while array effects and uncertainties in the particle size and shape distribution improve the agreement with experiment to some extent, the effect is too small to account for the deviation. To assess the contribution of the DFs, we also calculated the optical sensitivities using DFs from experiment. The results obtained using different experimentally measured DFs exhibit a spread that is comparable to the deviation between the optical sensitivities obtained using the calculated DFs and the direct measurements. This suggests that additional contributions to the DFs associated with scattering by, for example, defects are crucial. This highlights that the calculated DFs are in principle suitable for predictive simulations but that more work is required to assess more quantitatively the impact of samplespecific scattering and broadening channels.

Regardless, our simulations thus suggest that the optical sensitivity of individual nanoparticles cannot be significantly improved beyond the plateau that has been demonstrated here. For an array of nanoparticles, the optical sensitivity can, however, be improved by arranging them on a lattice and/or shifting the LSPR farther away from the avoided crossing, i.e., by increasing the nanodisk diameter far beyond 100 nm. In fact, the experimentally measured optical sensitivity in the farinfrared (i.e., for nanodisks with the largest diameters) clearly approaches the plateau value predicted here. Importantly, while from the experimental data one might anticipate the optical sensitivity to increase further for an even more extreme shift to the infrared (regardless of whether it is experimentally realizable or not), the present simulations provide strong evidence that there is an upper bound that is intrinsic for this kind of material and approach.

The actual sensitivity *S* comprises both the optical sensitivity \tilde{S} and a thermodynamic factor S_{sob} which depends on the solubility of hydrogen as a function of partial pressure. This thermodynamic sensitivity can vary rather strongly not only with the partial pressure of H₂ but the Au concentration. In particular for partial pressures near the α/β phase transition one can observe a strongly nonmonotonic variation with Au concentration with a maximum near the Au concentration at which the α/β two-phase region closes. While in experiments ensemble effects are likely to broaden this maximum, it remains a possible means to increase the actual sensitivity by alloying.

Another strategy to improve a H sensor is to increase the H uptake at a fixed hydrogen pressure, which means that for a specific optical sensitivity (measured optical shift per unit of absorbed H), lower hydrogen pressures can be detected. Here, we have shown that this can be achieved via an ordered L12-Pd₃Au phase, which can be obtained through annealing at high hydrogen pressures. Our simulations show that the optical response is largely unaffected by the presence of the ordered phase, which means that the optical sensitivity is unchanged. The H uptake, on the other hand, is significantly higher for the ordered phase which yields a detection limit 1-2 orders of magnitudes lower than for the disordered phase, well below the threshold stipulated by the U.S. Department of Energy (Figure 8). The possibilities provided by tuning the thermodynamic sensitivity through alloying demonstrated here have potentially notable implications for the field of plasmonic H sensing.

METHOD SECTION

Calculation of Dielectric Functions. SQSs²⁷ were generated with the ICET package⁴³ using the method and parameters suggested in ref 44. The ionic positions and cell

shapes of these structures were relaxed with density-functional theory (DFT) in the projector augmented wave formalism as implemented in the Vienna ab initio simulation package (version 5.4.1, PAW 2015)^{45,46} using the vdW-DF-cx exchange–correlation functional,⁴⁷ until residual forces were below 10 meV/Å and stresses below 1 kbar. In these calculations, the wave functions were expanded in a plane wave basis set with a cutoff of 384 eV, and the Brillouin zone was sampled with a Γ -centered grid with a *k*-point density corresponding to 19 × 19 × 19 *k*-points in the primitive cell. Occupations were set using the first-order Methfessel–Paxton scheme with a smearing parameter of 0.1 eV.

DFs were calculated using random phase approximation (RPA)-linear-response time-dependent density-functional theory as implemented in the GPAW package^{48,49} (version 19.8.1 with GLLB-sc patched for extended metallic systems). The macroscopic DF was calculated in reciprocal space through the linear density-density response function with wave functions expanded in a plane-wave basis with a cutoff at 350 eV. The BZ was sampled with a Γ -centered grid with a density corresponding to 61 *k*-points in each direction for the undecorated (monoelemental) primitive cell, and occupation numbers were smeared using a Fermi-Dirac distribution with a width of 0.1 eV.

DFs were obtained in the optical limit by evaluation at q = 0. As discrete *k*-point sampling precludes intraband transitions with $q \rightarrow 0$ and $\omega \rightarrow 0$, an intraband term, $\omega_{\rm p}^2/(\omega + i\eta)^2$, was added to the DFs, where $\omega_{\rm p}$ is the calculated plasma frequency, using the broadening parameter $\eta = 0.01$ eV. The ground state wave functions were obtained with the GLLB-sc exchange–correlation functional,⁵⁰ whereas dynamic exchange–correlation effects were neglected; i.e., the RPA approximation was used. In the optical limit, adiabatic local-density approximation (ALDA) results approach the RPA results, meaning virtually identical results would have been obtained if dynamic exchange–correlation effects had been taken into account using ALDA.

Finite-Difference Time-Domain Simulations. Single particle extinction spectra were calculated using finite-difference time-domain simulations as implemented in the MEEP software.⁵¹ The computational cell consists of a $Pd_{1-y}Au_yH_x$ nanodisk (represented by the calculated DFs fitted to a Lorentzian representation as described in Supplementary Note 1; refs 52-54) placed on a 100 nm thick SiO₂ substrate with constant refractive index 1.478. The nanodisk has truncatedcone geometry with fixed cone angle $\theta = 60^{\circ}$, varying (bottom) diameter d and height h (Figure 2a). To account for the expansion that occurs with H absorption in a real system, the system size is scaled with H content according to the lattice parameter expansion obtained from the DFT calculations. The source is a Gaussian pulse at normal incidence corresponding to the energy interval 1-5 eV. To mimic an infinite system, a 100 nm thick perfectly matched layer encloses the cell at a distance of at least 100 nm from the nanodisk in all directions, to enable a sufficiently large vacuum region. The cell is described in Cartesian coordinates with a grid resolution of 0.5 pixels/nm.

A detailed description of the procedure of calculating extinction spectra can be found in Supplementary Note 2. In the data analysis, a peak in the extinction spectrum is defined as a point in the extinction spectra with larger amplitude than the two neighboring points. For the peak shift sensitivity calculation, only peaks with higher amplitude than the mean over the entire spectrum were included. Calculating the sensitivity involves a linear fit of the studied shift with H

content. In this work, we employ Huber regression as implemented in SCIKIT-LEARN⁵⁵ for this purpose. T-Matrix Simulations. Nanodisk array extinction spectra were calculated using a T-matrix based approach as implemented in SMUTHI.56 The method allows for calculating the response of a substrate-supported single disk of varying geometry, as well as large arrays composed of hundreds of such particles. Note that for the nanodisk arrays, cylindrical disks where simulated instead of truncated cones, which has a negligible effect on the resulting extinction spectra (Figure S5). Arrays were treated either as amorphous and described statistically by the pair correlation function dependent on a minimum center-to-center distance between disks (as previously demonstrated^{57,58}) or as arrays consisting of 100 randomly distributed resonators with explicitly defined positions (see Figure S16b). For the latter case, size distribution within the array is also taken into account.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsanm.2c01189.

Additional visualizations of the DOS and DFs, nanodisk shape effects on the extinction spectrum, additional visualizations of extinction spectra, extended analysis of the double peak phenomenon of PdH, near-field enhancement obtained from electrodynamic simulations, hydrogen uptake as a function of the pressure and alloy composition, extended analysis of the peak shift sensitivity, effect of artificial broadening on the extinction spectra and the resulting sensitivity, particle size distribution and array effects on the optical response, optical response of the L1₂-ordered phase, and visual representation of the DF fitting procedure (PDF)

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Notes

The authors declare no competing financial interest. The raw dielectric functions are provided via Zenodo at https://doi.org/10.5281/zenodo.5833929.

ACKNOWLEDGMENTS

This work was funded by the Knut and Alice Wallenberg Foundation (Grants 2015.0055 and 2019.0140), the Swedish Foundation for Strategic Research Materials framework (Grant RMA15-0052), the Swedish National Research Council (Grants 2018-06482 and 2020-04935), and the Excellence Initiative Nano at Chalmers. T.P.R. acknowledges funding from Academy of Finland under Grant Agreement 332429. T.J.A. acknowledges support from the National Science Center, Poland, via Project 2019/35/B/ST5/02477. The computations were enabled by resources provided by the Swedish National Infrastructure for Computing (SNIC) at NSC, C3SE, and PDC partially funded by the Swedish Research Council through Grant Agreement 2018-05973 and the Interdisciplinary Center for Mathematical and Computational Modelling via Project GC84-51.

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(37) The definition of the onset is arbitrary and does not have to be based on the peak. Here we use 50% of the peak amplitude as a convenient measure when comparing sensitivity over the span of geometries.

(38) The exact concentration of the hydride is 67% for the Silkin et al. DF, 62% for the Palm et al. DF, and 58% for the Vargas et al. DF due to the concentrations of the available data.

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Supporting Information

Computational Design of Alloy Nanostructures for Optical Sensing of Hydrogen

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Supplementary Notes

Supplementary Note 1: Lorentzian representation of dielectric functions

To make the calculated dielectric functions (DFs) amenable to finite-difference time-domain (FDTD) simulations with the MEEP software (1), they need to be represented as a sum of N Lorentzian functions,

$$\epsilon(\omega) = 1 + \sum_{j=1}^{N} \frac{x_j \omega_j^2}{\omega_j^2 - \omega^2 - i\omega\gamma_j} = 1 + \sum_{j=1}^{N} x_j L_j(\omega)$$

$$\tag{1}$$

where x_j , ω_j and γ_j are parameters to be determined. As the computational cost has a direct dependence on the number of Lorentzians N, we want to keep N low. We have found the following strategy effective to determine N, x_j , ω_j and γ_j such that the calculated DF are accurately represented (for a graphical illustration of the procedure, see Figure S20):

- 1. Create a matrix with elements $A_{ij} = \operatorname{Re} L_j(\omega_i)$ and $A_{ij} = \operatorname{Im} L_j(\omega_i)$ in the upper and lower parts, respectively. This matrix will have twice as many rows as there are frequencies ω_i in the data obtained by linear-response time-dependent density-functional theory (LR-TDDFT) in the frequency range to be fitted (one for the real part and one for the imaginary part; in our case we use 100 values of ω_i in the photon energy range 0.7 to 10.7 eV for a total of 200 rows). Each column corresponds to a Lorentzian, and we use a 2D grid in the $\omega_j - \gamma_j$ plane with 200 values of ω_j in the range (0–10.7 eV) and 200 values of γ_j logarithmically spaced in the range 10^{-6} to 1×10^2 eV, for a total of 40,000 (ω_j, γ_j) pairs. The matrix A thus has dimensions $200 \times 40,000$.
- 2. With the thus obtained matrix we have an underdetermined system of equations $A_{ij}x_j = \epsilon(\omega_i)$, where $\epsilon(\omega_i)$ are as-calculated values of the DF at discrete frequencies ω_i . We solve this regression problem with the OMP (Orthogonal Matching Pursuit) algorithm as implemented in SCIKIT-LEARN (2) and wrapped in HIPHIVE (3). OMP allows us to control the sparseness of the solution, *i.e.*, the value of N. We test for values of N between 4 and 10 and choose the solution that has the lowest BIC (4) value.
- 3. Given the thus obtained solution with N oscillators, we fine-tune the 3N parameters $(x_j, \omega_j, \gamma_j)$ by solving the non-linear system of equations (1) in a least-squares sense (with the left-hand side given by the ascalculated LR-TDDFT values), as implemented in SCIPY (5).

Supplementary Note 2: Extinction spectra calculation with FDTD

During the FDTD simulations, Maxwell's equations are evolved in time until the intensity of the electric field at the disk center has decayed by a factor of at least 10^{-6} from its previous maximum value. The power $P_{in}(\omega)$ transmitted through a closed surface surrounding the nanodisk is calculated by integration of the Poynting vector. The absorption cross section σ_{abs} is then obtained by normalizing with the incoming intensity $I(\omega)$,

$$\sigma_{abs}(\omega) = \frac{P_{\rm in}(\omega)}{I(\omega)}.$$
(2)

The incoming intensity $I(\omega)$ is calculated from a preceding simulation without the nanodisk and correspond to the transmitted power through a surface above the nanodisk normalized by area. Calculation of the scattering cross section σ_{sca} requires subtracting the fields from the preceding substrate-only simulation before calculating the transmitted power $\tilde{P}_{out}(\omega)$ through a surface surrounding the nanodisk in the outward direction (as opposed to the inward direction for absorption),

$$\sigma_{sca}(\omega) = \frac{\tilde{P}_{\text{out}}(\omega)}{I(\omega)}.$$
(3)

Lastly, the extinction cross section $\sigma_{ext}(\omega)$ can be calculated as the sum of absorption and scattering,

$$\sigma_{ext}(\omega) = \sigma_{abs}(\omega) + \sigma_{sca}(\omega). \tag{4}$$

Supplementary Figures

Figure S1: Real part of dielectric function for gold content ranging from 0 to 45.8 % (a–l) and ranging from no hydrogen (blue) to full hydrogen loading (red).

Figure S2: Imaginary part of dielectric function for gold content ranging from 0 to 45.8 % (a–l) and ranging from no hydrogen (blue) to full hydrogen loading (red).

Figure S3: Electronic density of states at increasing Au content (a–i) in the range from no hydrogen (blue lines) to full hydrogen loading (red lines).

Figure S4: Comparison of our calculated dielectric functions (dashed, orange lines) to dielectric functions from the literature (blue lines). Panels (a,c,e) shows dielectric functions for pure Pd, panels (b,d,f) show the Pd hydride, with hydrogen content as reported by the respective authors and compared to dielectric functions as calculated in this work with the same hydrogen content (to within 0.5 percentage points). The data has been taken from Palm *et al.* (Ref. 6, a and b, measured on 200 nm films), Vargas *et al.* (Ref. 7, c and d, measured on 30 nm films), and Silkin *et al.* (Ref. 8, e and f, calculated with a method similar to ours).

Figure S5: Extinction spectra for nanodisks with slightly different shape. In addition to the truncated cone shape from the main study, we consider here a disk, a disk with a rounded upper edge (consisting of a half ellipsoid stacked on a disk) and a half ellipsoid (see inset for a side view). All of the disk have the same diameter and height (measured at the largest point). The degree of rounding/tapering of the upper disk edge does not have a significant effect on the spectra.

Figure S6: Absorption and scattering spectra for the same systems as in Figure 2.

Figure S7: Extinction spectra for PdH_x nanodisks with varying height h and AR. For the larger systems, we observe a single peak shifting to lower energies with increased H content, which is what is typically observed for localized surface plasmon resonance H sensors. For smaller systems, e.g. for $30 \text{ nm} \times 3$, a second peak appears with increased H content, shifting to higher energies. The peak splitting feature appear roughly over a diagonal in the figure. Above this imagined diagonal, we again observe a single peak but here it generally shifts to higher energies with increased H content, in contrast to the behavior for larger systems.

Figure S8: Peak amplitude (a) and position (b) for PdH nanodisks with varying height and AR and the corresponding dielectric function (c). With the extended energy interval, the connection between features in the DF and optical spectra is even more clear.

Figure S9: Extinction spectra and near fields at the resonances for PdH nanodisks with height 20 nm.

Figure S10: Hydrogen uptake in Pd–Au alloy as a function of Au concentration at pressures ranging from 0.1 mbar to 1 bar. Full lines show uptake when the Pd–Au sublattice is fully disordered, stars when the Pd–Au sublattice is ordered in the L1₂ configuration. These data are based on the model and simulations developed in Ref. 9). Orange diamonds show hydrogen uptake experimentally determined in Ref. 10 using quartz crystal microbalance. The discontinuity in hydrogen uptake, which is particularly wide at low Au concentration, is a consequence of the phase transition between the dilute α phase and the relatively dense β phase, which in pure Pd occurs just over 0.01 bar.

Figure S11: Thermodynamic sensitivities (a) $S_{\rm sol} = dc_{\rm H}/dp_{\rm H_2}$ and (b) $\tilde{S}_{\rm sol}$ using a reference pressure of $p_{\rm H_2}^{\rm ref} = 1 \times 10^{-9}$ bar at 300 K as a function of Au concentration at different H₂ partial pressures at a temperature of 300 K.

Figure S12: Optical peak shift sensitivity (a) and disk diameter (b) as a function of peak position at 0% H for varying Au concentration and disk heights 20, 30 and 40 nm (the marker size is proportional to height). The peak position follows a linear trend with respect to the disk diameter which is only slightly shifted with Au concentration or height.

Figure S13: Optical sensitivity (based on peak shift) for Pd nanodisks of varying size and the corresponding root-meroot mean squared error (RMSE) of the linear fit. The RMSE is larger right before the sensitivity reaches the plateau, due to the large effect from peak splitting.

Figure S14: H sensitivity based on the extinction shift at a certain wavelength/energy. The normalized extinction as a function of H concentration at varying energy (as indicated by the color scale) for the geometries in Figure 2 without Au (a–d) as well as the corresponding fitted slope (e–h) for each line.

Figure S15: Effect of size distribution on extinction spectra. Two different geometries (specified as $d \times h$) and compositions are considered, and the size distribution is introduced by varying the diameter uniformly in the interval $d \pm 10\%$ while keeping the particle volume fixed (blue lines). The difference between the single particle spectra at the average geometry (orange line) and the ensemble average over the size distribution (black line) is very small, indicating that a uniform size distribution is not enough to introduce broadening of the extinction spectrum.

Figure S16: (a) Extinction spectra of a single disk with height 30 nm and radius 50 nm placed on a substrate with refractive index n = 1.47813 (black line) and of amorphous arrays consisting of such disks. Amorphous arrays described statistically by the minimum center-to-center distance, cc (expressed in units of disk diameter) are marked with blue/cyan lines. Arrays of 100 randomly distributed disks with explicitly defined positions are marked with red lines, where σ corresponds to the deviation of disk radii drawn from a normal distribution with standard deviation equal to $r_0\sigma/3$ and the mean $r_0 = 50$ nm. Disk heights were adjusted to ensure constant volume. (b) An exemplary array of size-dispersed disks used in simulations (c) with a distribution of the disk radii. Surface densities for the amorphous arrays of cc = 1.63, cc = 2 and the explicit array used are approximately 26% and 17%, respectively. As expected, peak position and its width vary with the array density for the amorphous arrays. For explicitly defined arrays the resonance is blue-shifted and broader than the one corresponding to the amorphous array of the same surface density. Introducing size deviation does not drastically change the response, but results in a slight broadening of it, in agreement with (11). (d) The resulting optical peak shift sensitivities compared to the single particle and experimental sensitivities presented in the paper (in gray). Note that, except for the data in gray, the sensitivities here are calculated based on only the peaks at 0% and 67% H.

Figure S17: Optical peak sensitivity and dielectric functions when the interband transition (IBT) is removed from $PdH_{0.67}$ to mimic the case of extreme broadening of this feature. Note that, except for the data in gray, the sensitivities here are calculated based on only the peaks at 0% and 67% H which explains the rather large scatter. Without the interband transition, the optical sensitivity remains at high, positive values.

Figure S18: Effect of artificial Gaussian broadening of extinction spectra. The degree of broadening is determined by the width w of the Gaussian functions and w = 0 corresponds to no broadening. By applying broadening, the optical sensitivity essentially changes from a step function to a roughly linearly trend, similar to what is observed experimentally.

Figure S19: Extinction spectra and dielectric functions for the ordered $L1_2$ phase of $Pd_{75}Au_{25}$ compared to disordered special quasi-random structures of $Pd_{75}Au_{25}$ at varying H content. The spectra share the same general features, although the $L1_2$ structures have more fine structure. The fine structure would probably be averaged out in a real sample due to broadening, so we conclude that both cases should give similar measured optical response.

Figure S20: (a) Schematic overview of fitting of LR-TDDFT results to representation as sum of Lorentzians, as described in the Materials and Methods section of the main paper. A grid (ω_j, γ_j) of Lorentzian frequencies and broadening parameters (gray circles) is created (in practice it is much denser than shown here), and orthogonal matching pursuit is used to select a small number (4–10) of Lorentzians that reproduce the LR-TDDFT data well (green circles). These Lorentzians are then fine-tuned by solving the non-linear problem (orange circles). (b–c) The resulting functions (orange, dashed lines) match the original data (blue lines) very well over the energy interval used for FDTD simulations, here exemplified by the dielectric functions of Pd₀.75Au₀.25 with 0, 25, 50, 75, and 100% hydrogen.

Figure S21: Comparison between extinction spectra from the literature and our calculated spectra for various alloy and H concentrations. Our calculations are conducted as in the main text for truncated cones with dimensions as indicated in the figure titles $(d \times h)$. The has been taken from Rahm *et al.* (Ref. 12, upper row, measured experimentally and calculated using FDTD), Poyli *et al.* (Ref. 13, bottom row to the left, calculated using FDTD) and Langhammer *et al.* (Ref. 14, bottom row to the right, measured experimentally, the H concentrations are approximated since only the pressure is reported).

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