Gold–Rhodium Nanoflowers for the Plasmon-Enhanced Hydrogen Evolution Reaction under Visible Light

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ABSTRACT: State of the art electrocatalysts for the hydrogen evolution reaction (HER) are based on metal nanoparticles (NPs). It has been shown that the localized surface plasmon resonance (LSPR) excitation in plasmonic NPs can be harvested to accelerate a variety of molecular transformations. This enables the utilization of visible light as an energy input to enhance HER performances. However, most metals that are active toward the HER do not support LSPR excitation in the visible or near-IR ranges. We describe herein the synthesis of gold–rhodium core–shell nanoflowers (Au@Rh NFs) that are composed of a core made up of spherical Au NPs and shells containing Rh branches. The Au@Rh NFs were employed as a model system to probe how the LSPR excitation from Au NPs can lead to an enhancement in the HER performance for Rh. Our data demonstrate that the LSPR excitation at 533 nm (and 405 nm) leads to an improvement in the HER performance of Rh, which depends on the morphological features of the Au@Rh NFs, offering opportunities for optimization of the catalytic performance. Control experiments indicate that this improvement originates from the stronger interaction of Au@Rh NFs with H2O molecules at the surface, leading to an icelike configuration, which facilitated the HER under LSPR excitation.

KEYWORDS: gold, rhodium, nanoflowers, plasmonic catalysis, hydrogen evolution reaction

INTRODUCTION

Hydrogen (H2) represents one of the most promising energy carriers to address challenges related to renewable energy.1 H2 production from electrochemical water splitting has attracted massive interest in the past few years.2–4 Although electrochemical water splitting is straightforward in principle, it only represents 4% of the global H2 production due to its high cost and low efficiency.5–7 The water-splitting reaction encompasses H2 and O2 production through the hydrogen evolution reaction (HER) and oxygen evolution reaction (OER), respectively.6 In this context, noble-metal NPs have been successfully employed as electrocatalysts for H2 generation via the HER, in which Pt-based catalysts represent the state of the art.7–9 Nevertheless, due to the associated high costs and low abundance of noble metals there is a strong drive for further improving their activity.10

Strategies to improve the performance of noble-metal NPs toward the HER involve size and shape control, bimetallic compositions, and a wealth of hybrid systems.11–13 It has been recently established that the localized surface plasmon resonance (LSPR) excitation in plasmonic NPs can be harnessed to accelerate and drive a variety of chemical and electrochemical reactions.14–17 When plasmonic NPs support-
enables the marrying of plasmonic and catalytic properties, in which the plasmonic material acts as an antenna harvesting energy from light through the LSPR excitation and the catalytic component uses this energy to drive or accelerate a transformation of interest.\textsuperscript{10,41} Thus, it becomes clear that this approach can also be beneficial in the context of improving HER performances and enabling renewable energy input (sunlight) and low overpotentials.

We report on the controlled synthesis of gold–rhodium core–shell nanoflowers (Au@Rh NFs) that are comprised of a spherical Au nanoparticle (NP) core and a shell containing Rh branches or small NPs. This material was employed as a model system to probe how the LSPR excitation from the Au NPs can lead to an enhancement in the catalytic activity of the Rh in the shells toward the HER. While this approach has been demonstrated on several plasmonic–catalytic combinations, examples containing Rh and exploring the bimetallic plasmon-enhanced catalytic activity remain limited.\textsuperscript{42–46} Nevertheless, Rh has important catalytic properties toward a wealth of transformations that would benefit from plasmonic enhancement and control.\textsuperscript{47,48} Although the synthesis of Au@Rh core–shell NPs has been reported\textsuperscript{44–46} applications have focused on the HER and not on the effect of plasmonic excitation on HER performances, indicating the need of a deeper understanding of the HER under light irradiation. Here, by employing the HER as a proof of concept transformation, it was demonstrated that the LSPR excitation at 533 nm (and 405 nm) led to an improvement in the HER performance. Our control experiments indicate that this improvement originated from the stronger interaction of Au@Rh NPs with H\textsubscript{2}O molecules at the surface, leading to an icelike configuration, which facilitated the HER under LSPR excitation. The catalytic behavior was found to be dependent on the morphological and compositional features of the Au@Rh NFs, offering future opportunities for optimizing the performance by tuning the morphology, structure, and composition.

\section*{Experimental Section}

Materials and Methods. All of the reagents were used without further purification: tetrachloroauric(III) acid trihydrate (HAuCl\textsubscript{4}·3H\textsubscript{2}O, 99\% Sigma-Aldrich), rhodium chloride (RhCl\textsubscript{3}, 98\%, Sigma-Aldrich), hexadecyltrimethylammonium bromide (CTAB, C\textsubscript{16}H\textsubscript{33}N\textsubscript{3}O·3H\textsubscript{2}O, 99\% Sigma-Aldrich), sodium citrate trihydrate (C\textsubscript{6}H\textsubscript{5}Na\textsubscript{3}O\textsubscript{7}·3H\textsubscript{2}O, 99\% Sigma-Aldrich), L-ascorbic acid (C\textsubscript{6}H\textsubscript{8}O\textsubscript{6}, ≥99\% BioXtra, Sigma-Aldrich), sodium iodide (NaI, 99.5\% Sigma-Aldrich), sodium bromide (NaBr, 99\% Synth), sodium chloride (NaCl, 99\% Synth), polyvinylpyrrolidone (PVP, Sigma-Aldrich, MW 10000 g mol\textsuperscript{−1}), tetrabutylammonium bromide (TBAB, C\textsubscript{16}H\textsubscript{35}NBrN, 98\% Sigma-Aldrich), sodium borohydride (NaBH\textsubscript{4}, 98\% Sigma-Aldrich), hydroquinone (C\textsubscript{6}H\textsubscript{4}O\textsubscript{2}, 99.5\%, Sigma-Aldrich), Nafion 5\% (C\textsubscript{2}F\textsubscript{5}O\textsubscript{2}S·C\textsubscript{2}F\textsubscript{2}O\textsubscript{2}, 15–20\%, Sigma-Aldrich), 2-propanol (C\textsubscript{3}H\textsubscript{6}O\textsubscript{2}, 99.9\% hypergrade for LC-MS LiChrosolv, Merck), and Carbon Vulcan XC-72R. All aqueous solutions were prepared with Millipore water (18.2 M\textsubscript{Ω}).

**Instrumentation.** High-resolution transmission electron microscopy (HRTEM) images and initial energy dispersive X-ray (EDX) spectra were taken on a JEOL JEM 2100 instrument operating at an accelerating voltage of 200 kV. The catalysts were redispersed in isopropanol by centrifugation and ultrasound bath agitation and then drop-cast onto transmission electron microscopy (TEM) grids (Formvar/Carbon Film coated, 200 mesh, Cu) and dried under ambient conditions. A JEOL ARM300F instrument was used to acquire high-angle annular dark field (HAADF) scanning transmission electron microscope (STEM) images and EDX spectrum images at a 300 kV accelerating voltage. EDX data were acquired using an Oxford Instruments XMAX 100 detector. HAADF-STEM images were acquired with an inner detector semicircle of 77 mm rad. UV–vis spectra were recorded on a Shimadzu UV-2600 spectrophotometer. The samples were suspended in water and placed in quartz cuvettes with an optical path of 1 cm. The metallic content of the suspension and deposited materials was measured by inductively coupled plasma atomic emission spectroscopy (ICP-OES) on a Spectro Arcos instrument.

**Synthesis of Gold Nanoparticles (Au NPs).** First, 30 mL of distilled water and 300 μL of AuCl\textsubscript{4}·(aq) (25 mmol L\textsuperscript{−1}) were placed in a round-bottom flask and the flask was placed on a heating plate with stirring at 130 °C. When the mixture was boiling, 900 μL of a sodium citrate solution (38.7 mmol L\textsuperscript{−1} or 1%) was added and the reaction proceeded for 10 min. Finally, the red suspension obtained was washed by successive rounds of centrifugation and removal of the supernatant and then concentrated to 4 mmol L\textsuperscript{−1} (concentration given in terms of Au content). The suspension concentration was determined by inductively coupled plasma atomic emission spectroscopy (ICP-OES). The as-produced Au NPs had a diameter of 14.8 ± 1 nm as measured by TEM (Figure S1a).

**Synthesis of Rhodium Nanoparticles (Rh NPs) as Controls for the Electrochemical Studies.** A 3.70 mg portion of NaBH\textsubscript{4}, 9.3 mg of RhCl\textsubscript{3}, and 87 mg of PVP 10000 were weighed and transferred into a milling jar. The milling experiments were conducted with a 14 mL PTFE milling jar using three ZrO\textsubscript{2} balls (10 mm, 3.17 g each), giving a ball to powder mass ratio (bpr) of 95:1. The milling device consists of a vibrating ball mill with a fixed vibration frequency of 27.5 Hz (SL-38, SOLAB-Brazil). The Rh nanoparticles were obtained after 1 h of milling and had a diameter of 2 nm as determined by TEM (Figure S1b).

**Synthesis of Gold–Rhodium Nanoflowers (Au@Rh NFs).** A 0.1 g portion of CTAB, TBAB, or PVP was placed in a round-bottom flask containing 18.6 mL of distilled water. This mixture was sonicated for 5 min and heated to 90 °C with stirring. After 5 min, 1 mL of the previously prepared Au NP suspension (4 mmol L\textsuperscript{−1}), 50 μL of an aqueous NaI, NaBr, or NaCl solution (0.01 mol L\textsuperscript{−1}), and a specific volume (150, 350, 550, or 750 μL) of a RhCl\textsubscript{3} aqueous solution (20 mmol L\textsuperscript{−1}) were added to the reaction mixture. The amount of Rh was varied to investigate how the NF morphology and the plasmonic–catalytic properties changed as a function of the Rh content. After 5 min, 1 mL of ascorbic acid, NaBH\textsubscript{4}, or hydroquinone solution (0.04 mol L\textsuperscript{−1}) was added, and the reaction was allowed to proceed for 30 min. The brownish black solution obtained (Figure S2) was washed by successive rounds of centrifugation and removal of the supernatant and deposited on a Carbon Vulcan support by wet impregnation for further electrochemical measurements. After deposition on the Carbon Vulcan support, all samples were analyzed by ICP-OES and the metal loading in all cases was around 2%.

**HER Experiments.** Electrochemical testing was performed with an electrochemical cell consisting of a reversible hydrogen electrode (RHE) reference and a Pt wire as the auxiliary electrode. The working electrode was a glassy-carbon-disk electrode (d = 1.5 mm) modified by drop-casting 10 μL of
catalyst ink (5 mg of the catalyst dispersed in 2 mL of water) and 5 μL of a Nafion 0.05% solution. The electrolyte was 1 mol L\(^{-1}\) KOH solution, and the cell was N\(_2\)-saturated to avoid the formation of reactive oxygen species. A hydrodynamic condition of 1600 rpm controlled by a Pine AFMSRCE RDE apparatus was used to avoid accumulation of hydrogen bubbles on the electrode surface. To investigate the catalyst HER performance, linear potential, triangular potential, and potential step perturbations were executed and controlled by a PGSTAT302N Autolab potentiostat/galvanostat. For the light-driven experiments, modular lasers of 405, 533, and 650 nm of 300, 200, and 390 mW cm\(^{-2}\) power density, respectively, were positioned 5 cm from the cell. The laser was turned on 5 min before starting the measurement, to avoid power fluctuation. The perturbation programs were the same as those under dark conditions. The electrochemically active surface area (ECSA) was estimated from the electrochemical double-layer capacitance from each catalyst. Cyclic voltamograms were recorded at several scan rates (10–1000 mV s\(^{-1}\)) from 0 to 0.6 V\(_{\text{RHE}}\). The double-layer capacitance (\(C_\text{dl}\)) was calculated from the slope of the current (at 0.30 V\(_{\text{RHE}}\)) versus scan rate linear plots. The \(C_\text{dl}\) values of the catalysts were divided by the \(C_\text{dl}\) value of pure Carbon Vulcan, as the support also makes a contribution to the surface area.

**In Situ Fourier Transform Infrared (FTIR) Spectroscopy.** An electrochemical thin-layer cell with a CaF\(_2\) window was used in the spectroelectrochemical experiments, using a reversible hydrogen electrode (RHE) as a reference electrode and Au foil surrounding the working electrode as an auxiliary electrode. The working electrode was a shiny Au disk (\(d = 5\) mm), modified by the same procedure described in the electrochemical methods. The cell was continuously purged with N\(_2\), and the electrolyte used was 1 mol L\(^{-1}\) KOH. The experiments were performed in a reflection absorption setup (IRRAS) with a Nicolet 6700 spectrometer (MCT-B detector, gain of 8, 4 cm\(^{-1}\) resolution, 64 accumulations), and the potential perturbation was controlled by a PGSTAT302N Autolab potentiostat/galvanostat. The open-circuit potential (OCP) was measured for 5 min, and potential step perturbations were applied from the OCP to \(-0.24\) V\(_{\text{RHE}}\) during each spectrum acquisition.

**X-ray Photoelectron Spectroscopy (XPS) Experiments.** The X-ray photoelectron spectroscopy measurements were performed with a Specs FlexPS system equipped with a Phoibos 150 2D CCD detector and a XR50 X-ray source with
a nonmonochromatic Al Kα (1486.61 eV) source. The catalyst ink was drop-casted over a Si substrate, entirely covering the surface, as no Si signal (the main transition for SiO₂ is Si2p at 103.3 eV)³⁹ was detected. All spectra were registered in a high-vacuum chamber at pressures below 10⁻⁹ mbar and were corrected by the C₁s signal (284.5 eV).⁴⁹,⁵⁰

### RESULTS AND DISCUSSION

**Synthesis of Au@Rh NFs and LSPR Enhanced HER Performances.** The synthesis of gold–rhodium nanoflowers (Au@Rh NFs) was performed by a seeded growth approach using preformed Au NPs as physical templates for the deposition of different Rh amounts on their surface. Figure 1a–c shows TEM images for the Au@Rh NFs obtained by the addition of 150, 350, and 550 μL of 20 mmol L⁻¹ RhCl₃, respectively, to the reaction mixture. The Rh atomic percentages in the Au@Rh NFs was determined by EDX (Figure 1d) and correspond to 10 (Figure 1a), 18 (Figure 1b), and 32 at. % (Figure 1c). These samples were therefore denoted as Au₉₀Rh₁₀ (Figure 1a), Au₈₂Rh₁₈ (Figure 1b), and Au₆₈Rh₃₂ (Figure 1c). It can be observed that all of the samples were relatively monodisperse, and the size of the NFs corresponded to around 19.3 ± 2, 22.9 ± 2, and 23.8 ± 1 nm for samples Au₀₉₀Rh₁₀ (Figure 1a), Au₈₂Rh₁₈ (Figure 1b), and Au₆₈Rh₃₂ (Figure 1c), respectively. Histograms of the size distribution are shown in Figure S3. The increase in size correlates with the increased amount of RhCl₃ added to the synthesis and the higher Rh content in the NPs, indicating the size increase can be ascribed to an increased Rh deposition at the Au surface.

This is confirmed by HRTEM images for Au₉₀Rh₁₀ (Figure 1d), Au₈₂Rh₁₈ (Figure 1e) and Au₆₈Rh₃₂ (Figure 1f) NFs. All images show Au cores (~15 nm diameter) covered in small Rh NPs (~2 nm in diameter), with the Au and Rh identifiable by their lattice fringe spacings (e.g., Figure 1h) or using STEM-EDS elemental mapping (Figure 1j). Figure 1d reveals that in the Au₉₀Rh₁₀ NFs Rh is present as individual NP islands or single-crystal branches on the Au surface. At higher Rh loadings (Au₈₂Rh₁₈ and Au₆₈Rh₃₂ in Figure 1f and Figure 1e, respectively), an increasingly high number of Rh particles or branches covered the Au NPs, yielding an Au@Rh core–shell morphology for the NFs at higher Rh content. Although each individual island/branch appears to be single-crystalline, the tortuous morphology, as illustrated in Figure 1g for Au₆₈Rh₃₂, suggests this is formed by an oriented attachment of separate Rh particles. It can be observed that each branch/island at the surface is <5 nm in diameter and displays lengths of around 5 nm. When the Rh content in the NFs was further increased, it became relatively monodisperse, and the presence of individual Rh NPs was detected in addition to the Au@Rh NFs. This result indicates that, as the Au surface becomes completely covered with Rh, no further Rh deposition at the surface of the Au@Rh can take place and the additional Rh NPs remain in solution.

Figure 2 shows the UV–vis extinction spectra registered from an aqueous suspension containing the Au NP seeds and the Au₉₀Rh₁₀, Au₈₂Rh₁₈, and Au₆₈Rh₃₂ NFs. The Au seed NPs display a band centered at 522 nm that is assigned to the LSPR dipolar mode.⁵¹ The intensity of this band decreased with the deposition of Rh on the surface in Au₉₀Rh₁₀ NFs, as would be expected due to the suppression of the Au LSPR extinction with the deposition of a nonplasmonic metal at its surface. For Au₈₂Rh₁₈ and Au₆₈Rh₃₂ NFs, in addition to a further decrease in the intensity of this band, a red shift to 548 nm and broadening of the signal are observed. This can be assigned to the change in dielectric constant due to the increased Rh deposition at the surface. As the overall NF size is also increased, scattering can also contribute to the broadening of the signal. Importantly, all NFs display extinction signals characteristic of the Au LSPR in the visible range, despite the Rh deposition at the Au surface.⁵² Additionally, an increase in the intensity in the short-wavelength region from 350 to 400 nm is detected, which scaled with the Rh content and is assigned to the increasing Rh LSPR in the UV range.⁵³,⁵⁴ The signal in this region can also have contributions of Au interband transitions below 515 nm.

To further investigate the formation mechanism for the Au@Rh NFs, we performed control experiments to unravel the effect of I⁻ ions on the resulting NF morphology. In the absence of I⁻ ions, the TEM images (Figure S5a) show a core–shell structure, consisting of the Au NP seed surrounded by a lower-density, uniform shell layer containing the Rh NP seeds but no Rh branches. This result shows that the presence of I⁻ ions is essential to enable the formation of Rh-based branches at the surface of the Au NP seeds. Here, it is plausible that I⁻ ions can act as a stabilizer of the Rh surface facets that favor the formation of the branched morphology via oriented attachment. This was further confirmed by replacing I⁻ ions by Br⁻ ions in the synthesis (Figure S5b,c, respectively). While Br⁻ ions also promoted the deposition of Rh islands or branches at the Au NP surface, their morphology was not as uniform and well-defined as in the case of I⁻ ions. As the interaction of Rh with halides is expected to increase in the order I⁻ > Br⁻ > Cl⁻,⁵⁵ it is plausible that I⁻ would act as a better stabilizer relative to the other halides for Rh surface facets that favor the deposition of Rh branches and lead to the morphology of the Au@Rh NFs.

Control experiments replacing CTAB by TBAB (Figure S6a) and PVP (Figure S6b) did not lead to the formation of the NFs. This indicates that both CTAB and I⁻ ions are essential to the synthesis of Au@Rh NFs. The effect of the reducing agent was also investigated by replacing ascorbic acid with NaBH₄ (Figure S6c) and hydroquinone (Figure S6d). In both cases, the Au@Rh NF morphology was not detected, illustrating that the use of a mild reducing agent coupled with...
the use of CTAB and I\textsubscript{aq} ions was imperative to the formation of Au@Rh NFs. Although iodide and bromide are known for poising surfaces in catalytic applications, the presence of iodide was not detected by XPS (Figure S7). Although some bromide was detected and could influence the activity (decrease HER performance), it was detected in all samples, indicating that the chemical environment and, thus, the effect would be the same in all cases. This indicates that further optimization of the washing procedure can lead to improvements in the HER activity.

Table S1 summarizes all the experimental conditions that were investigated for the synthesis optimization of Au@Rh NFs. Finally, we monitored the morphological changes and growth evolution as a function of time by stopping the synthesis at different time intervals (5, 10, 15, and 30 min) and analyzing the generated NPs by HRTEM, as shown in Figure S8 for the sample composition Au\textsubscript{68}Rh\textsubscript{32}. As the synthesis time varied from 5 to 30 min, a gradual increase in the deposition of Rh small islands or branches at the Au NP surface was detected, together with the formation of individual Rh NPs after 30 min. These results indicate that, in addition to the amount of Rh precursor employed during the synthesis, the synthesis time can be employed to control the degree of Rh deposition at the surface of Au NPs.

Having verified the morphology produced during the synthesis, we then investigated the Au@Rh NFs as model systems to consider how plasmonic effects can influence the materials performance toward the HER. The hypothesis is that the optical/plasmonic properties of the Au core can enhance the Rh HER performance under visible light illumination via plasmonic electrocatalysis. For the electrochemical experiments the nanostructures were impregnated in Carbon Vulcan to increase the material conductivity. The metallic loading (Au + Rh) was determined by ICP-OES to be around 2% in all cases, which for Au\textsubscript{68}Rh\textsubscript{32} corresponds to 0.06 μg of Rh and 0.14 μg of Au. We performed the experiments at pH 14 to avoid corrosion and improve the stability of the catalyst.

We started by recording triangular potential perturbations at 50 mV s\textsuperscript{−1} from 0 to 1.2 V\textsubscript{RHE} (illustrated schematically in Figure S9a) to evaluate the electrochemical profile for each material, as presented in Figure S9. Peaks between 0 and 0.2 V\textsubscript{RHE} were attributed to the H\textsubscript{2} adsorption–desorption reaction (H\textsubscript{adps}) on Rh\textsuperscript{56,57} and are noticeable in all three samples tested (Au\textsubscript{90}Rh\textsubscript{10}, Au\textsubscript{82}Rh\textsubscript{18}, and Au\textsubscript{68}Rh\textsubscript{32}). Rh shows a stronger adsorption of hydrogen in comparison to Au\textsuperscript{58}. Thus, as the Rh ratio increases, so does the H\textsubscript{adps}.

Polarization curves at 5 mV s\textsuperscript{−1} from 0.1 to −0.3 V\textsubscript{RHE} under hydrodynamic conditions were recorded with and without light illumination at the working electrode for Au\textsubscript{90}Rh\textsubscript{10}, Au\textsubscript{82}Rh\textsubscript{18}, and Au\textsubscript{68}Rh\textsubscript{32} NFs, as shown in Figure 3a. Monometallic Au and Rh seed NPs were also evaluated as reference materials. A green laser (533 nm, 200 mW cm\textsuperscript{−2}) was used as the LSPR excitation source, which was chosen due to its proximity to the extinction band of the NFs (Figure 2). It can be observed that the monometallic Au and Rh NPs do not show any difference in activity in the applied potential range for light and dark conditions. An increase in the HER activity is observed as the Rh content of the Au@Rh NFs increases, under both dark and light conditions (Figure 3a,b). The overpotential at 10 mA cm\textsuperscript{−2} ECSA (η\textsubscript{10}) under dark conditions corresponds to 0.25, 0.20, and 0.17 V\textsubscript{RHE} for Au\textsubscript{90}Rh\textsubscript{10}, Au\textsubscript{82}Rh\textsubscript{18}, and Au\textsubscript{68}Rh\textsubscript{32} NFs, respectively. These data indicate that it is only possible to produce hydrogen in the probed potential range under our experimental conditions when both metals are coupled in the bimetallic NFs. The LSPR excitation produced by light illumination led to an enhancement in the HER activity in the Au\textsubscript{82}Rh\textsubscript{18} and Au\textsubscript{68}Rh\textsubscript{32} NFs, decreasing the η\textsubscript{10} values by 40 and 30 mV\textsubscript{RHE}, respectively (Figure 3c). This corresponds to a decrease in η\textsubscript{10} values of 20 and 18%, respectively. Surprisingly, no changes in the η\textsubscript{10} value were detected for the Au\textsubscript{90}Rh\textsubscript{10} NFs upon illumination. As the HER studies were
performed under the same loading of metal (Au + Rh), and the performance and light enhancement are dependent on the amount of Rh, it is plausible that the electrode must hold a minimum critical concentration of Rh so that the performance becomes high enough to be sensitive to the LSPR excitation. Under our experimental conditions, our results indicate that Au@Rh NF catalysts showed a better performance than Au(111) and commercial Rh/C in alkaline media. However, it is important to note that a more precise comparison would require normalization to the electrochemically active surface area (ECSA) and other experimental conditions, such as laser power density. Moreover, the Au68Rh32 performance was compared to those of other reported plasmonic catalysts with comparable overpotentials and increase in activity under light excitation. Table S2 presents a summary of the HER performances of various materials in alkali and acid electrolytes. It is noteworthy that plasmonic electrocatalysts for the HER are usually reported in acidic media and a higher overpotential is detected at higher pH. Nonetheless, acidic media show some drawbacks that limits their further application, such as corrosion and poor material stability.

We also investigated the dependence of the HER performance on excitation wavelength by employing laser irradiation at 405 and 650 nm (300 and 200 mW cm\(^{-2}\), respectively), as shown in Figure 3c. Neither of these wavelengths is matched to the main Au LSPR extinction band of the NFs (548 nm). As was observed for 533 nm excitation conditions, the Au68Rh32 NFs were found to be insensitive to light illumination. For Au82Rh18 and Au68Rh32 NFs, no significant changes in the overpotential were detected under 650 nm excitation compared to dark conditions, as expected. However, under 405 nm irradiation, a decrease in the overpotential was detected for Au82Rh18 and Au68Rh32, although it was a smaller decrease in comparison to that at 533 nm excitation. This enhancement in HER activity may be due to the partial contribution from the LSPR excitation tail of Rh at 405 nm (band maximum in the UV) and interband transitions in Au, which is reflected in the enhanced extinction seen in the UV–vis spectra (Figure 2). With regard to the interband transitions, they are observed for Au NPs in the high-energy regime due to direct d-to-s electronic excitation. It has been reported for some systems that the higher potential of these transitions can lead to higher photocatalytic activities relative to the LSPR (intraband) excitation, as hot electrons can be generated at the s band and holes can be generated in the d band. In this case, more hot carriers could generated from direct interband excitation transitions relative to intraband excitations, which may lead to higher photocatalytic activities relative to LSPR excitation. Therefore, a means to optimize plasmonic-catalytic performances can be obtained by leveraging these transitions.

To further confirm that the enhancement of the HER was due to LSPR excitation (at 530 nm) and interband transitions (at 405 nm) in the NFs, the on–off j–t transients were recorded under chopped light illumination at both 533 (Figure 4a) and 405 nm (Figure 4b) for the Au68Rh10, Au82Rh18, and Au68Rh32 NFs. All transients were registered at −0.1 V\(_{\text{RHE}}\) which is in the Tafel region. It can be observed that all samples present a fast and reproducible current response to on–off illumination cycles. The steady currents under light illumination were wavelength-dependent, being greatest for the 533 nm excitation. By subtraction of the steady current before (\(j_{\text{dark}}\)) and after (\(j_{\text{light}}\)) the illumination, the photocurrents could be calculated as 0.15, 0.27, and 0.30 mA cm\(^{-2}\) ECSA for Au68Rh10, Au82Rh18, and Au68Rh32, respectively, indicating an increase in the photocurrent with the Rh loading.
without light excitation were obtained from the quasi-steady-state polarization curves, as shown in Figure 4c. The HER mechanism in alkaline medium can be described by the Volmer, Heyrovsky, and Tafel steps, as shown in eqs 1–3. The values in parentheses refer to the Tafel slope, which comes from the Tafel equation ($\eta = a \log j$), where $j$ is the current density and $a$ is the Tafel slope.

Volmer step: $\text{H}_2\text{O} + e^- \rightarrow \text{H}_{\text{ads}} + \text{OH}^-$ (118.2 mV dec$^{-1}$)  

(1)

Heyrovsky step: $\text{H}_2\text{O} + \text{H}_{\text{ads}} + e^- \rightarrow \text{H}_2 + \text{OH}^-$ (39.4 mV dec$^{-1}$)  

(2)

Tafel step: $\text{H}_{\text{ads}} + \text{H}_{\text{ads}} \rightarrow \text{H}_2$ (29.6 mV dec$^{-1}$)  

(3)

These steps represent competitive processes of adsorption/desorption of hydrogen atoms occurring at the catalyst surface. All steps play a key role necessary to obtain excellent HER performance. Without light illumination, the average Tafel slope corresponded to 89 ± 4 mV dec$^{-1}$. This indicates that the mechanism cannot be explained by a single rate-limiting step (Heyrovsky or Volmer) without further kinetic studies. Intriguingly, there were no significant changes in the Tafel slopes under light illumination, indicating that the HER mechanism remains unchanged under plasmonic excitation, in agreement with previous studies. These results therefore indicate that the LSPR excitation can lead to enhancements in the HER activity in the Au@Rh NFs, but the HER mechanism remains unchanged.

Investigating the Role of the Bimetallic NF Composition over the Water Adsorption Interactions (under Dark Conditions). It is well-known that HER activity presents a linear relationship with the electrode working function. This relation is due to the generation of adsorbed hydrogen, as presented in the Volmer step and corroborated by DFT simulations. Even though it is in good agreement with acidic electrolytes, the main problem with this kind of assumption was that for all these investigations high vacuum was needed for the experimental setup or assumed for the simulations. When these proprieties were investigated by a consideration of more realistic environmental conditions, fine-tuning was possible. A volcano plot was obtained in which several proprieties were considered such as the work needed for a proton transfer from the outer Helmholtz plane (OHP) to the inner plane (IHP), the desolvation of this proton in the IHP that occurs during the adsorption process, and the charge transfer energy between the metal and the proton for the generation of this adsorbate. In this, Au is present in the weak adsorption branch, presenting a very low activity, while Rh is close to the volcano top but is already sitting at the stronger adsorption branch. This study has also shown that the increase in the work function not only affects the adsorption energy but also influences in the pre-exponential factor of the Arrhenius equation for the HER. This model, however, does not consider in detail the electrolyte–electrode interactions and fails to describe how different materials behave at higher pH values, since under alkaline conditions, the reactants should be the protons obtained by the water self-ionization or the water molecules themselves, as shown in eq 1. Since water should now be considered, it is also important to highlight that the water adsorption conformation affects the way the O–H bond will be more or less easily broken during the Volmer step. The electrochemical activity dependence on the water adsorption conformation is not new; actually, it was already proposed by Trasatti for the HER, and it has been shown in the literature that it can affect the electrochemical activity for different reactions. Therefore, it is proposed that the water structure orientation at the double-layer region will give insights into investigations on the HER mechanism.

Next, we wanted to investigate how the bimetallic composition of the NFs (without light excitation) influence their activities toward the HER. In this case, we were interested in probing if the bimetallic composition could enable different adsorption interactions with water that could lead to improvements in the HER relative to the monometallic NPs. We performed in situ Fourier transform infrared spectroelectrochemistry experiments to gain further insights into the intermediates involved in the hydrogen evolution reaction. These experiments were performed for Au$_{68}$Rh$_{32}$ NFs and Au and Rh NPs, aiming at further understanding the importance of the bimetallic compositions to the improved HER performances in the NFs. The in situ FTIR spectra were recorded under potential step perturbation. Figure 5a shows the spectra obtained for Au$_{68}$Rh$_{32}$ NFs as a function of the applied potential (Figure S10a,b shows the Au and Rh NPs spectral counterparts, respectively). At approximately 1225 and
1154 cm$^{-1}$ two sharp bands are observed, which were assigned to the asymmetric ($\nu_{as}$) and symmetric ($\nu_s$) stretching of CF$_2$ from Nafton. The decrease in the CF$_2$ stretching modes with the potential suggests water incorporation into Nafton.$^8$ The bands detected at all potentials at approximately 2920 and 2850 cm$^{-1}$ were attributed to the asymmetric and symmetric CH$_2$ stretching modes of an amine from the CTAB.$^8$ The bending mode of the adsorbed water ($\delta$(HOH))$^8$ was observed at 1618, 1664, and 1650 cm$^{-1}$ in Au, Rh, and Au$_{68}$Rh$_{32}$, respectively. A lower frequency of $\delta$(HOH) indicates a stronger water adsorption, as the increase in charge transfer from the oxygen lone pairs decreases the band

Figure 6. Stark tuning plots and absorbance versus potential plots for deconvoluted bands in the range (a, b) 3240−3320 cm$^{-1}$, (c, d) 3400−3530 cm$^{-1}$, and (e, f) 3530−3650 cm$^{-1}$ for Au, Rh, and Au$_{68}$Rh$_{32}$. 
frequency. Thus, the Au$_{68}$Rh$_{32}$ NPs showed a stronger water adsorption in comparison to Rh, indicating that combining both metals in the nanostructure can improve the catalyst interaction with H$_2$O. Nonetheless, a direct comparison cannot be made between the bimetallic NPs and Au NPs, as Au presents a weaker hydrogen bonding strength, leading the $\delta$(HOH) band to a lower frequency.\(^{58,83}\)

The band centered at $\sim$1700 cm$^{-1}$ for Au and Rh NPs (Figure S10a,b) was assigned to the bending mode of the adsorbed hydronium cation ($\delta$(H$_2$O$^+$)). More specifically, this corresponds to the interfacial hydronium–water complexes, such as H$_3$O$^+$, H$_2$O$_2^+$, and H$_2$O$_3^+$.\(^{84}\) Although the hydronium complex is not usually observed at higher pH due to its low concentration,\(^{85}\) it can be locally produced during the HER due to the water dissociation process in the electrode at high pH.\(^{86}\) As the IR spectrometer in an IRRAS setup has a small optical path and poor mass transport, it can facilitate the observation of reaction intermediates even at low concentrations. This band appeared at higher wavenumber in pure Rh NPs (1720 cm$^{-1}$), indicating a stronger interaction in comparison to Au NPs (1690 cm$^{-1}$). This bending mode was not detected for Au$_{68}$Rh$_{32}$ NPs; it could be obscured by the $\delta$(HOH) band. In Au and Rh NPs, the $\delta$(H$_2$O$^+$) and $\delta$(HOH) bands are symmetrical and well-defined. On the other hand, the $\delta$(HOH) band in Au$_{68}$Rh$_{32}$ had a more asymmetric profile, with a tail toward higher frequencies at higher potentials. This could indicate the presence of more than one band. Nonetheless, as the intensity of this band was low, a deconvolution was not possible.

Moreover, a large band in the $\sim$3100–3800 cm$^{-1}$ region was attributed to the OH stretching mode ($\nu$(OH)) in Figure S5a. This band was deconvoluted into three bands centered at $\sim$3272, 3512, and 3637 cm$^{-1}$, with the fitting for Au$_{68}$Rh$_{32}$ NPs at $-0.12$ V$_{RHE}$ being shown in Figure S5b. Stark tuning (wavenumber versus potential) and absorbance versus potential plots are shown in Figures 6.

The Stark plots for Au NPs were constant for all three bands, and no shift in wavenumber was observed in the probe potential range. This indicates that the interaction of Au with adsorbed water is weak for all conformations. The first deconvoluted band centered at 3249, 3268, and 3289 cm$^{-1}$ (Figure 6a) for Au, Rh, and Au$_{68}$Rh$_{32}$, respectively, is attributed to an icelike water conformation (Figure S11b). This water conformation has a strongly bonded hydrogen, due to the Fermi resonance of the $\delta$(HOH) binary overtone and the $\nu$(OH).\(^{87}\) The estimated slopes from the Stark tuning plot for this band (Figure 6a) were $-270$ and $-570$ cm$^{-1}$ V$^{-1}$ for Rh and Au$_{68}$Rh$_{32}$, respectively. Here, the higher the slope, the stronger the interaction between the metal and hydrogen, suggesting adsorption and, consequently, the existence of this water conformation in the IHP. Therefore, these data suggest that the presence of both Au and Rh in the bimetallic NPs led to an even stronger interaction with water in comparison to both individual Au and Rh NPs. This band increased in intensity with an increased applied potential (Figure 6b), the Au absorbance variation being smaller than that for the other catalysts as a result of the weaker H$_2$O interactions.\(^{88,89}\) The stronger interaction of an icelike water structure with the bimetallic NF could explain its higher catalytic activity in comparison to monometallic Au and Rh, as the icelike conformation leads to more aligned and oriented water molecules that act as sources of protons due to enhanced water autoprotolysis.\(^{86,89}\) In this way, more H$_{ad}$ is formed and the Volmer step, which is the initial step and probably one of the limiting steps toward the HER, can be accelerated. In summary, the icelike water conformation could contribute to affecting the electron-transfer chemistry because, as there is more H$_{ad}$ in the catalyst surface, the hot electrons could be injected in the adsorbed water molecules and lead to the cleavage of O–H. If there is more H$_{ad}$ this step is then facilitated.

The second deconvoluted band around 3400 cm$^{-1}$ (Figure 6c,d) is assigned to hydrogen-bonded water in the bulk,\(^{87,90}\) in which water molecules assume a more flat-lying structure (Figure S11a). These bands were centered at 3423, 3482, and 3528 cm$^{-1}$ for Au, Rh, and Au$_{68}$Rh$_{32}$, respectively, suggesting a stronger interaction of the water with Au$_{68}$Rh$_{32}$ in comparison to individual Au and Rh NPs separately, due to its higher wavenumber. Once more, this band did not shift for Au NPs over the applied potential range, whereas Rh and Au$_{68}$Rh$_{32}$ showed a shift to higher wavenumbers with the potential. As can be seen in Figure 6c, Au$_{68}$Rh$_{32}$ also showed a higher slope in the Stark tuning plot (440 cm$^{-1}$ V$^{-1}$) in comparison to the Au and Rh NPs (0 and 375 cm$^{-1}$ V$^{-1}$, respectively), likely as a result of the stronger interaction. Moreover, the absorbance for this band (Figure 6d) also increased with potential as a result of these surface interactions. It was also suggested in the literature\(^{74}\) that water at the third layer (already OHP) already presents a bulk conformation. In this way, the cases in which no Stark effect was observed, this species is probably not interacting directly with the electrode.

Finally, the third deconvoluted bands (Figure 6e,f) were centered at 3537, 3628, and 3637 cm$^{-1}$ for Au NPs, Rh NPs, and Au$_{68}$Rh$_{32}$ NPs, respectively. These bands correspond to water in the solvation shell of the potassium ions in the electrolyte (Figure S11).\(^{76,82,91}\) Au and Au$_{68}$Rh$_{32}$ showed a similar behavior, with small variations for this band (Figure 6e), indicating that the adsorption of solvated K$^+$ ions is weaker for these catalysts. Nonetheless, Rh NPs showed the highest slope in the Stark plots and wavenumber plots (Figure 6f), indicating that the interaction of rhodium with the solvated complex is much stronger in pure Rh NPs. The presence of solvated cations on the electrode can hinder the HER by a steric effect, explaining the higher activity in Au$_{68}$Rh$_{32}$ in comparison to Rh NPs.

These results indicate that the bimetallic composition enables a stronger interaction with water (stronger water adsorption), leading to higher activities relative to the monometallic counterparts. The observation of a preferential water conformation (icelike structure) in the IHP for the most active catalyst indicates that the electron transfer should be facilitated by its presence. It has been proposed in the literature\(^{74}\) that in alkaline media electron transfer and water self-ionization are simultaneous steps for the Volmer mechanism. The data presented here help with an understanding of the chemical environment and corroborate the hypothesis presented in the literature about the water conformation and electrode–electrolyte interaction influence on Faradaic processes. However, since dynamic spectr electrochemical measurements were not possible due to the setup limitations, quantitative data could not be achieved.

Moreover, the light excitation can contribute to further enhance the performance of the bimetallic NPs due to the plasmonic effect. If we compare the activities between the bimetallic NPs and the monometallic samples, our data suggest that the bimetallic composition plays a major role in the
improvements of performances relative to the LSPR effect, as the increase in activity from the monometallic samples to the bimetallic NFs in the dark is more significant than the additional enhancement provided by the plasmonic excitation.

On the basis of these results, Figure 7 schematically suggests a mechanism for the enhanced HER performances based on the bimetallic composition and plasmonic effects in the Au@Rh NFs. The bimetallic composition contributes to increasing the strength of adsorbed H$_2$O at the surface of the NFs, which preferentially leads to an icelike or interfacial water conformation, as suggested by the in situ FTIR. This enables a significant increase in HER performance relative to the monometallic systems, showing that this conformation is favoring the electron transfer simultaneously with the water self-ionization, even in the absence of light excitation. Under 533 nm light irradiation, hot electrons (and holes) are generated from the LSPR excitation in Au. These hot electrons can flow to the Rh surface, where they can contribute to the activation of adsorbed H$_2$O species, leading to a decrease in the overpotential.

Figure 7. Schematic representation of electron transfer pathways that lead to an enhancement in the HER performances following the LSPR excitation in the Au@Rh NFs. Under 533 nm excitation, hot electrons generated from the Au LSPR excitation can flow to the Rh surface, where they can contribute to the activation of adsorbed H$_2$O species, leading to a decrease in the overpotential.

CONCLUSION

We have described herein a facile method for the synthesis of core–shell Au@Rh (antenna@reactor) NFs comprised of spherical Au nanoparticles (NPs) and shells containing Rh islands/branches (smaller than 5 nm). The synthesis was performed by seeded growth, and the degree of Rh deposition in the NFs could be tuned by adjusting the amount of Rh precursor employed during the deposition step or controlling the synthesis time. The Au@Rh NFs were employed as a model system to investigate the effect of the LSPR excitation from the Au NPs and their enhancement toward the HER of the Rh shells as a function of the NF composition. Our results demonstrated that the LSPR excitation at 533 nm led to an improvement in the HER performances. Specifically, NFs containing 68 Au at. % and 32 Rh at. % (Au$_{68}$Rh$_{32}$) displayed the highest performance toward the HER and plasmonically enhanced HER. Experiments for lower and higher excitation wavelengths confirmed the role of the LSPR excitation toward the HER and suggest that the increase in the HER for the bimetallic NFs relative to the monometallic counterparts was due to a change in the interaction with adsorbed water molecules. In situ FTIR spectra suggested that adsorbed water species assumed an icelike configuration in the bimetallic NFs, thus facilitating the HER. The results presented demonstrate that, although Rh only supports LSPR in the UV region, plasmonic effects due to visible light irradiation can be harvested to enhance the catalytic activities of Rh, by the synthesis of hybrid nanomaterials containing Rh coupled with Au. We believe that the results presented herein provide insights into the physical and chemical parameters that need to be controlled to optimize the HER performance in Rh NPs via plasmonic excitation in the visible range and may inspire the design and development of Rh-based nanomaterials toward this and other plasmonically enhanced catalytic applications. Possible future directions include other NP designs such as bimetallic systems that enable decreased Rh contents in the form of alloys or single-atom alloys (containing Rh as the single-atom components at the Au surface) and an under-
standing of why Au@Rh NPs lead to a stronger interaction with water.

■ ASSOCIATED CONTENT

* Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscatal.1c02938.

Information on the characterization of the Au, Rh, and Au@Rh NPs by TEM, XPS, and cyclic voltammetry, study of the electrocatalytic performance by in situ FTIR spectra, performance comparison with other reported HER electrocatalysts, modeling of optical properties, and models for water absorption (PDF)

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Notes
The authors declare no competing financial interest.

■ ACKNOWLEDGMENTS

The authors thank the Brazilian agencies CNPq and the São Paulo Research Foundation FAPESP (2015/26308-7) for financial support. M.P.d.S.R. also thanks the FAPESP for the fellowship granted (2018/16846-0). P.H.C.C. thanks the Jane and Aatos Erkko Foundation, the University of Helsinki, HELSUS, and the Academy of Finland for financial support. The authors thank Diamond Light Source for access and support in use of the electron Physical Science Imaging Centre (Instrument E01 and proposal number NT26559) that contributed to the results presented here.

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