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4. Enhancement of hydrogen evolution rates on platinum electrodes by controlling interfacial water reorganization
ABSTRACT

The hydrogen evolution on platinum electrodes is a milestone reaction in electrocatalysis as well as an important reaction towards sustainable energy storage. Although the hydrogen evolution mechanism has been the subject of numerous studies, the pH dependent kinetics of this reaction is not yet fully understood. We present here a detailed kinetic study of the hydrogen adsorption and evolution reaction on Pt(111) in a wide pH range. Impedance measurements show that the hydrogen adsorption and hydrogen evolution are both slow in alkaline media, which is consistent with the observation of a shift in the rate-determining step for H₂ evolution with increasing pH value. Adding nickel to the Pt(111) surface lowers the barrier for the hydrogen adsorption rate and thereby enhances the hydrogen evolution rate. These observations are explained by a new model which highlights the role of interfacial water reorganization to accommodate charge transfer through the electric double layer, the energetics of which is controlled by how strongly water interacts with interfacial field. In alkaline media, the potential of zero charge is very positive compared the hydrogen evolution potential, leading to a higher activation energy for reorganizing water. Nickel lowers the potential of zero charge of Pt(111), as confirmed by laser-induced temperature jump experiments, and thereby lowers the barrier for hydrogen adsorption. Our findings and model shed new light on the origin of the slow kinetics for the hydrogen evolution reaction in alkaline media, solving a long-known problem toward efficient hydrogen production.

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4.1. Introduction

There is a global call for industrial processes that combine economic progress with long-term preservation of natural resources. In terms of technological advances for sustainable energy production, there is a recent renewed interest to realize the so-called hydrogen economy by photocatalytic water splitting or by the combination of photovoltaics with water electrolysis. Methane steam-reforming is currently the most cost-efficient technology available for hydrogen production, but unsustainable in the long run as it is still based on the deployment of fossil fuels. In order for the hydrogen economy to meet our future energy demands, there are, however, various fundamental bottlenecks to be overcome, such as that related to the efficient catalysis of the associated multi-proton multi-electron transfer reactions. Essentially, significant advances are required to lower the high inherent costs of electrocatalysts, by increasing the efficiency of water oxidation, by the replacement of scarce and expensive catalyst materials by earth-abundant alternatives, and by maximizing their durability and lifetime. Substantial efforts have thus been devoted to lowering the costs of the electrodes necessary for water splitting. Recent theoretical works exposed the mechanistic features of the oxygen evolution reaction (OER), taking place at the anodes of electrolyzers, in which water is oxidized to produce molecular oxygen and protons. These insights motivated several reports on earth-abundant and highly-efficient OER anode materials that typically work at high pH. Unfortunately, the price to be paid for the use of alkaline conditions is a significant overpotential at the cathode, where the hydrogen evolution reaction (HER) takes place.

It has long been known in the electrochemistry literature that the kinetics of both the hydrogen evolution reaction (HER) and hydrogen oxidation reaction (HOR) on platinum are significantly slower in alkaline media than in acidic media. The elucidation of the molecular-level origin of this problem would be of obvious importance for the further development of alkaline electrolyzers and alkaline fuel cells. Marković et al. have recently shown how the oxophylicity of the interface, as
modified by adsorbing a small amount of Ni(OH)$_2$ on Pt(111), may improve the kinetics of the HER/HOR, and ascribed this effect to the favorable interaction of surface adsorbed OH$_{\text{ads}}$ with the relevant intermediates. Mechanistic studies on the HER$^{10a-c, 11}$ have traditionally correlated reaction rates with thermodynamic descriptors, in particular the strength of the bond between hydrogen and the metal electrode, following the Sabatier principle$^{12}$. Gasteiger $et$ $al.$ and Yan $et$ $al.$ have suggested$^{11c, 13}$ that a pH-dependent H-binding energy lies at the origin of the pH-dependent HER/HOR kinetics, and therefore they concluded that the H-binding energy is and remains the sole descriptor for the HER/HOR reaction. Specifically, Yan $et$ $al.$ have considered the pH-dependent shifts of the voltammetric peaks in the so-called “underpotential deposition” (UPD) hydrogen region of polycrystalline platinum as evidence for such a pH-dependent H-binding energy. However, the molecular-level origin of the pH-dependent H-binding energy has remained elusive. More importantly, we have recently argued$^{14}$ that the nature of the “hydrogen” peaks on polycrystalline platinum are unlikely to be associated with the adsorption of hydrogen only, but also include the effect of the adsorption of oxygenated species on (110) and (100) sites. Therefore, their peak potentials are not unambiguous indicators of H-binding energy. Moreover, on a Pt(111) electrode, there is no significant shift of the H-UPD with pH, but there is still a very significant pH-dependence of the HER/HOR kinetics (see below). Therefore, it appears that in spite of the undeniable success of the traditional models, they overlook important kinetic details, and that a consistent explanation of why HER/HOR on Pt is slow in alkaline media is still missing.

In this work, we present a detailed kinetic study of the hydrogen adsorption and evolution reaction on a Pt(111) single-crystal electrode, in a wide pH range, and in the absence and presence of a small amount of Ni(OH)$_2$ promoter. Electrochemical impedance spectroscopy is used to accurately measure the charge transfer kinetics of hydrogen adsorption. We present a new model for the rate of hydrogen adsorption step, based on the idea that the barrier for this reaction depends on how close or remote the electrode potential is in relation to the potential of zero
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The laser-induced temperature jump technique is then used to measure the potential of maximum entropy (pme), which is closely related to the pzc, to show that at pH 13, the addition of Ni(OH)$_2$ to Pt(111) indeed shifts the pme/pzc closer to the hydrogen adsorption region. We attribute the impact of the pzc on the activation barrier of hydrogen adsorption to the energy penalty associated with the reorganization of interfacial water to accommodate charge movement through the double layer. In acid media, the pzc/pme of Pt(111) is close to the hydrogen region, and the energy of reorganization of the interfacial water to move a proton through the double layer is relatively small. In alkaline media, the pzc/pme of Pt(111) is far from the hydrogen region (i.e. close to the OH$_{ads}$ region) and the corresponding strong electric field existing at the electrode/electrolyte interface in the hydrogen region leads to a large reorganization energy for interfacial water when OH$^-$ transfers through the double layer. In general terms, our results show how a cost-effective, earth-abundant metal as nickel, in the form of nickel hydroxide, promotes the reorganization of water networks at the electrode-electrolyte interface by shifting the pzc closer to the equilibrium potential of the HER reaction, thereby enhancing the reaction rate for the hydrogen evolution at high pH values. Our model also suggests a new strategy for designing new and better electrocatalysts in aqueous media, highlighting the role of the interfacial solvent reorganization on ion transfer steps.

4.2. Experimental Details

Experiments were carried out in a fluorinated ethylene propylene (FEP, Nalgene) electrochemical cell for alkaline solutions, whereas a glass cell was preferred for acidic solutions, using a three-electrode assembly at room temperature in both cases. The cells and glassware were initially cleaned by boiling in a 1:1 mixture of concentrated sulfuric acid and nitric acid, rinsed thoroughly and boiled with ultra clean water (Milipore® MiliQ; resistivity >18.2 MΩ.cm) five times
before each set of experiments. All the glassware and cells were kept in a permanganate/sulfuric acid mixture when not in use.

Solutions of different pH value were prepared as follows: for pH 1 we used solutions of 0.1 M HClO$_4$ (Merck 70%, Suprapur). In the preparation of pH 1.8 we used a mixture of 0.01 M HClO$_4$ + 0.090 M NaClO$_4$ (monohydrate, EMSURE), and for pH 2.5 the mixture was 0.001M HClO$_4$ + 0.099 M NaClO$_4$. Phosphate buffer solutions of 0.1 M were prepared for pH 4.3, 7, 9.1 and 9.8, using sodium dihydrogen phosphate and/or disodium hydrogen phosphate (Merck 99.99%, Suprapur) and ortho-phosphoric acid (Merck 85%, Suprapur) as indicated by the Henderson-Hasselbach equation for buffers. For pH 11 and 12 we prepared solutions of 0.001 M NaOH (Sigma-Aldrich, 99.99% trace metal basis) + 0.090 M NaClO$_4$, and 0.01M NaOH + 0.090 M NaClO$_4$ respectively. Solutions of 0.1 M NaOH were prepared for pH 13.

Bead-type single crystal platinum electrodes with a (111) orientation (<0.5 degrees) were used as working electrodes (WE). Before each experiment the electrode was flame-annealed for 1 min and cooled down to room temperature in a 3:1 proportion Ar + H$_2$ mixture (Linde 6.0). The electrode was transferred to the electrochemical cell with a protective droplet of deoxygenated water. All measurements were carried out with the Pt single crystal electrode in a hanging meniscus configuration. A platinum wire was used as counter electrode (CE). For the EIS measurements a home-made, self-contained hydrogen electrode (SCHE) was placed in the same electrolyte, with a Teflon connection in order to avoid contact between the solution and the glass. A 10 µF capacitor was connected, as a noise filter, between the RE and a platinum wire immersed in the solution. All potentials noted are referred to the reversible hydrogen electrode (RHE).

The argon was bubbled through a 3 M KOH trap before it entered the cell, in order to remove impurities which may be present from the tubing. The cell was placed inside a Faraday cage. Cyclic voltammetries and impedance spectra were
collected using a computer-controlled Ivium A06075 potentiostat (Iviumstat). Impedance spectra were measured with frequencies ranging from 10 kHz to 0.1 Hz and a peak-to-peak amplitude of 5 mV. The data was fitted to the equivalent electric circuit (EEC) shown in 2a, using a Python script, employing the Levenberg-Marquardt algorithm for non-linear least square regression.

For the deposition of a submonolayer of Ni(OH)$_2$ on the Pt(111) electrode we used a solution of 0.1M NaClO$_4$ and 5mM Ni(NO$_3$)$_2$ (Sigma-Aldrich 99.999%, trace metals basis). The solution was purged with Argon for 10 minutes and the Pt(111) electrode was modified via electroless instantaneous deposition, by placing the electrode in the deaerated solution for 10 s. After deposition, the Pt(111)-Ni electrode was rinsed with ultrapure water and brought to hanging meniscus configuration in a cell containing a 0.1 M NaOH solution, in order to perform cyclic voltammetry (3 cycles). The laser induced temperature jump experiment was performed as described elsewhere$^{15}$. Pulses of 5 ns of the second harmonic of a Nd-YAG laser (532 nm) were used as laser source, with an energy density around 8 mJ.cm$^{-2}$ (the laser beam diameter is 4 mm and the energy of the laser pulse is 1mJ), a value of energy small enough to prevent the damage of the Ni(OH)$_2$ deposit on the Pt(111) electrode. The laser energy was measured with a pyroelectric sensor head (Model M-935-10). The only effect from laser irradiation is the increase of the temperature of the interface.

4.3. Results and Discussion

Figure 1 illustrates the key observation that we wish to explain in this work, comparing the cyclic voltammeteries for a Pt(111) electrode at pH 11 (blue line), pH 12 (red line), pH 13 (turquoise line) and pH 1 (black, dashed line). All the measurements were collected using a scan rate of 50 mV.s$^{-1}$. The voltammetric curves are very similar in current and shape for the alkaline pH, showing the typical features for the Pt(111) surface orientation: the hydrogen adsorption region, known
as H-UPD (0.1 – 0.35 V_RHE), which remains almost unaltered for the several pH values presented (versus the RHE), the double layer region (0.35 – 0.55 V_RHE), and the hydroxyl adsorption-desorption region (0.6 – 0.8 V_RHE). The reduction current registered between 0.05 V_RHE and -0.1 V_RHE corresponds to the hydrogen evolution. We observe that the onset potential for the HER in alkaline solutions exhibits a shift towards negative potentials as the pH value increases. Hydrogen evolution clearly has the lowest overpotential at pH 1, even though there is only a small shift in the H-UPD region in comparison to alkaline media. We conclude from Figure 1 that a pH-dependent shift in the H-binding energy (if any) cannot explain the significant pH dependence of the HER overpotential on Pt(111).

Figure 1. Cyclic voltammetries for Pt(111) in 0.1 M perchloric acid (pH 1); 0.001 M NaOH + 0.099 M NaClO_4 (pH 11); 0.01 M NaOH + 0.09 M NaClO_4 (pH 12); 0.1 M NaOH (pH 13). Scan rate: 50 mV.s^{-1}. 
To demonstrate that the pH also affects the rate of hydrogen adsorption in the H-UPD region, we studied the kinetics of hydrogen adsorption in the 0.1-0.3 V potential window. Figure 2 summarizes the measurements made by means of electrochemical impedance spectroscopy as a function of potential and pH. The experimental admittance spectra were fitted using the equivalent electric circuit (EEC\textsuperscript{16}) presented in Figure 2a. This EEC corresponds to a simple heterogeneous adsorption step, forming an adsorbed intermediate without diffusion limitation. The charge transfer resistance, $R_{\text{ct}}$, is inversely proportional to the rate of the corresponding hydrogen adsorption reaction, i.e. $\text{H}^+ + e^- \rightarrow \text{H}_{\text{ads}}$ in acidic media, and $\text{H}_2\text{O} + e^- \rightarrow \text{H}_{\text{ads}} + \text{OH}^-$ in alkaline media. $R_{\text{sol}}$ stands for the resistance of the solution, whereas $C_{\text{DL}}$ and $C_{\text{AD}}$ represent the capacitance of the double layer and the pseudo-capacitance of the adsorbed hydrogen, respectively. Figure 2b shows the admittance plots, measured at 0.2 V\textsubscript{RHE} in solutions of 0.1 M ionic strength at different pH values. The data collected for the entire H-UPD region were fitted to the EEC shown in Figure 2a and can be found in Figure B1 in the Appendix B. From the Nyquist plots presented in Figure 2b we notice that for alkaline pH we observe two semicircles. The first semicircle at low frequencies (left side of each diagram) is related to the charge transfer resistance. For acidic media, only a single semicircle in the admittance plane is observed (not shown, see ref. 26), meaning that the H-UPD charge transfer process is too fast to be measured in acidic media\textsuperscript{16}. Therefore, not only the HER is slower in alkaline media, but also the H-UPD adsorption is slower, even if the thermodynamic driving force for hydrogen adsorption is the same (as all potentials are referred to the RHE scale and there is no observable pH dependence of the H-binding energy).

The $R_{\text{ct}}$ measurements were performed for solutions with different pH value for the H-UPD region and are shown in Figure 2c. The $R_{\text{ct}}$ for pH 7 is shown for comparison with the alkaline pH values (pH 7 was measured in a phosphate buffer, but since there is no phosphate adsorption in the H-UPD region\textsuperscript{17}, we do not expect an impact of phosphate on the hydrogen adsorption rate). From the figure we observe that the charge transfer is both pH- and potential-dependent. The
reaction becomes slower for higher pH values, pointing out that there is a reaction barrier for hydrogen adsorption that increases with pH. This effect cannot be described in terms of a presumed pH-dependent H-binding energy and must therefore have another origin.
Figure 2. a) Equivalent electric circuit for H-UPD region, featuring the double layer capacitance, $C_{DL}$, the capacitance associated to the hydrogen adsorption, $C_{AD}$, the charge-transfer resistance for hydrogen adsorption, $R_{CT}$, and the solution resistance, $R_{SOL}$. b)
Admittance Nyquist plots for Pt(111) in 0.1 M solutions for different pH values, measured
with frequencies ranging from 10 kHz to 0.1 Hz and an amplitude of 5 mV, at 0.2 V_RHE: the
dots represent the experimental data points collected, while the solid lines correspond to the
fit obtained using the EEC. The arrows point out how the components from the EEC in Fig.
2a are related to each semi-circle. The data for the entire H-UPD region can be found in
Figure A1, in the Appendix B. c) Inverse of the charge transfer resistance as a function of
the pH value. The R_CT was obtained from the fitting parameters for the admittance plots,
collected for Pt(111) in 0.1 M solutions with different pH value. The different colors represent
the potential at which the data was collected (see legend).

The mechanism for the hydrogen evolution at each pH value can be derived
from the relationship between the current density and overpotential, known as Tafel
plots. In order to estimate the Tafel slopes, we recorded polarization curves at 10
mV.s^{-1} under argon atmosphere, using 0.1 M solutions with pH values of 1, 1.8,
2.5, 4.3, 7, 9.1, 9.8, 11, 12 and 13, respectively. The polarization curves for these
solutions are shown in Figure B2 in the Appendix B. Figure 3 shows the Tafel
slopes as a function of pH, revealing the tendency of the slope to increase with pH,
as reported previously^{9b, 18} for pre-activated polycrystalline platinum electrodes.
The significant conclusion from the combination of Figures 1 and 2 is that not only
the rate of HER on Pt(111) changes with pH, but also the mechanism.

The relation of the rate-determining step in the mechanism to Tafel slope
analysis has been described elsewhere^{19}. Experimental reports^{18b, 20} on
polycrystalline and low-index single-crystal platinum electrodes under
hydrodynamic conditions, exhibit Tafel slopes of ca. 30 mV.dec^{-1} for acid solutions
and ca. 120 mV.dec^{-1} for alkaline solutions. Previous reported values of the Tafel
slope for the HER on Pt(111) in acidic media are 31 mV.dec^{-1} in perchloric acid by
Seto et al.^{20} and 35 mV.dec^{-1} in sulphuric acid by Kita et al.^{21} The interpretation of
the measured Tafel slope is as follows. A value around 120 mV.dec^{-1} implies that
the first electron transfer step (ET) is rate-determining (RDS), meaning that in
alkaline media the so-called Volmer step H_2O + e^- → H_ads + OH^- is the slowest
reaction step. A Tafel slope of ca. 40-30 mV.dec^{-1} implies that either the second ET
step is the RDS or the RDS is a chemical step preceded by two ETs, indicating that in acidic media the mechanism involves the Volmer step $\text{H}^+ + e^- \leftrightarrow \text{H}_{\text{ads}}$ in equilibrium, followed by a rate-determining Tafel step $2\text{H}_{\text{ads}} \rightarrow \text{H}_2$ or a rate-determining Heyrovsky step $\text{H}^+ + e^- + \text{H}_{\text{ads}} \rightarrow \text{H}_2$. These mechanistic implications are consistent with the observation that in alkaline solutions the hydrogen adsorption reaction is a slow reaction, whereas in acidic media this step is fast and in equilibrium.

![Tafel slopes measured for the hydrogen evolution reaction in 0.1 M solutions, in a wide pH range: 1, 1.8, 2.5, 4, 7, 9.1, 9.8, 11, 12 and 13 respectively. The description of the solutions used can be found in the experimental section.](image)

**Figure 3.** Tafel slopes measured for the hydrogen evolution reaction in 0.1 M solutions, in a wide pH range: 1, 1.8, 2.5, 4, 7, 9.1, 9.8, 11, 12 and 13 respectively. The description of the solutions used can be found in the experimental section.

Figure 4 compares the rate of hydrogen adsorption, as evaluated by EIS, and the rate of hydrogen evolution at pH 13 on bare Pt(111) and on Pt(111) in presence of a small amount of Ni(OH)$_2$. The Tafel plots for the HER (Fig. 4a) and
the inverse of the $R_{ct}$ for H-UPD (Fig. 4b) for Pt(111)/Ni(OH)$_2$ show a clear enhancement of both rates in alkaline media, as compared to Pt(111). The HER rate enhancement was already reported in previous works$^{10b,10d}$ by Marković et al.; the novelty here is that this enhancement also applies to the rate of H-UPD formation. From the Tafel plots in Figure 4a we can observe that the rate for hydrogen evolution in the presence of Ni(OH)$_2$ is now in between that for pH 13 and pH 1 on pure Pt(111), with a corresponding change in Tafel slope. This would suggest that the role of Ni(OH)$_2$ is to lower the barrier for the hydrogen adsorption reaction, rather than to change the energetics of the hydrogen intermediate or the mechanism.

![Figure 4](image-url)

**Figure 4.** a) Tafel plots for HER on for Pt(111) at pH 1 and pH 13 and on a Pt(111) electrode decorated with 0.1 ML of Ni(OH)$_2$ in 0.1 M NaOH (pH 13), under argon atmosphere. The polarization curves were registered at a scan rate of 10 mV.s$^{-1}$. b) The inverse of the charge transfer resistance as a function of potential for Pt(111) (black squares) and Pt(111) electrode decorated with ~0.14 ML of Ni(OH)$_2$ in 0.1 M NaOH, pH 13 (blue squares); argon atmosphere. The $R_{ct}$ was obtained from the fitting parameters for admittance plots (see Figure B3 in the Appendix B).
Our suggestion here is that the rate of hydrogen adsorption (and hence of hydrogen evolution if hydrogen adsorption is rate-determining) has a pH dependent barrier because the interfacial water structure at the Pt(111) electrode is pH dependent. During the hydrogen adsorption reaction, charge has to move through the interfacial double layer (H⁺ in acid and OH⁻ in alkaline). The rate of such a charge transfer reaction depends on the extent to which the solvation environment (i.e. the water) can accommodate this charge migration. If the interfacial water is easily reorganized, charge transfer through the double layer will be rapid; if the interfacial water is rigid and difficult to reorganize, charge transfer through the double layer will be slow. An important parameter to influence the extent to which the interfacial water can be reorganized is the interfacial electric field. The latter is determined by the charge separation between the metal electrode and the electrolyte solution, which in turn, is dependent on the position of the pzc. In the presence of adsorption reactions, distinction should be done between total and free charges. While the latter represents the true charge determining the electric field at the interface, the total charge, which includes also the charge stored in the bond of adsorbed species, is the only available parameter from electrochemical measurements. To obtain the true pzc (potential of zero free charge, pzfc), additional measurements and assumptions are necessary to separate adsorption reactions from the true capacitive processes. In the absence of a strong interfacial electric field, i.e. close to the potential of zero (free) charge (pzc), water is relatively free to reorient and hence easy to reorganize. In the presence of a strong interfacial electric field, i.e. further away from the pzc, water is much more rigid and hence more difficult to reorganize. As has been shown by previous work, while the pzfc cannot be directly measured, an estimation can be obtained from values of CO displaced charge. This is achieved by extrapolating the charge in the double layer region either into the hydrogen adsorption region (acid solutions) or into the hydroxyl adsorption region (alkaline solutions). In this way, a relatively constant value of pzfc is obtained, independent of pH in the SHE scale. This means that pzfc changes from the double layer region (0.34 V_RHE) in acidic
solutions to the onset of the surface oxidation (ca. 1.0 V\textsubscript{RHE}) in alkaline solutions\textsuperscript{23}. As a consequence, at a given potential on the RHE scale, the strength of the electric field and the energy associated with the (re)organization of interfacial solvent molecules will depend on pH. Since the H-UPD region is far from the pzc of Pt(111) in alkaline media, the hydrogen adsorption reaction is slow in alkaline solutions. In this model, a way to compensate this strong electric field is to use surface modifiers, such as nickel, that bring the pzc closer to the equilibrium potential of the hydrogen evolution reaction.

To assess whether such a shift in pzc and the associated energy of reorganization of the interfacial water could indeed explain the promoting effect of Ni(OH)\textsubscript{2}, we have employed the laser-induced temperature jump method. This technique uses short laser pulses to increase the temperature of the solvent at the interface for a very brief period of time\textsuperscript{25}. During the relaxation at open circuit, the temporal change in potential induced by the temperature perturbation is registered. The sign of the transient gives information about the average orientation of the interfacial solvent that existed at the potential before the potentiostatic control was removed and the short laser pulse was applied. In this interpretation, it is implied that adsorption processes are not fast enough to respond to the temperature perturbation. Specifically, the potential for which the polarity of the transient changes is identified as the potential of maximum entropy (pme), which is expected to be closely related to the pzc. Previous comparisons\textsuperscript{25b} between the pme and pzc of Pt(111) in acidic solution indeed show a close correspondence between the two quantities.

Details regarding the laser-induced temperature jump method can be found in the literature\textsuperscript{25b}. The method was used to determine the pme of the Pt(111) interface at pH 13, as well as the effect of a submonolayer amount of Ni(OH)\textsubscript{2} deposited on the Pt(111) surface on the pme. The results of the measurements are shown in Figure 5. We present the cyclic voltammetries for the bare and decorated electrode in Figure 5a. The transients for Pt(111) at pH 13 in Figure 5b are
negative at 0.5 V\textsubscript{RHE}. At that potential, the interfacial water molecules are oriented with their hydrogens pointing towards the electrode surface. We observe a change in polarity of the interfacial species at 0.675 V\textsubscript{RHE}. Assuming that the hydroxyl adsorption is not fast enough to contribute to the potential transient, this change in polarity suggests that from this potential the water molecules have an average orientation with the hydrogens pointing away from the electrode surface. The potential of maximum entropy, is therefore located close to 0.675 V\textsubscript{RHE} (some 25 mV lower than the pzc) at potentials above which the average orientation of the hydrogens in the water molecules are pointing away from the electrode surface (see Fig. 5). In this case, the difference between the pme and the extrapolated value of the pzfc\textsuperscript{23} would be a consequence of the hydroxyl adsorption on the water orientation and the double layer capacitance. What is clear is that pme and pzfc are both located above 0.6 V\textsubscript{RHE}, i.e., well above the value in acid solutions.

The laser-induced coulstatic potential transients collected for the Pt(111) electrode decorated with \textasciitilde~0.2 ML of Ni(OH)\textsubscript{2}, at pH 13, are shown in Figure 5c. Negative transients are observed for 0.60 V\textsubscript{RHE}. From 0.645 V\textsubscript{RHE} we observe a change in polarity of the transient, suggesting that the pme has shifted ca. 30 mV negatively with respect to the bare Pt(111). This shift in pme/pzc is in accordance with our model for the enhancement of the H-UPD and hydrogen evolution by the presence of Ni(OH)\textsubscript{2} on the surface as proposed above, relating the hydrogen adsorption and hydrogen evolution rates to the energy required for the water reorganization in the electrode interface. While the interpretation of the t-jump experiment is subject to some uncertainty regarding the interference of hydroxyl adsorption to the measured potential transient, previous results with other adatoms clearly demonstrated that the presence of electropositive adatoms at the electrode interface decrease significantly the value of the pme, and therefore the pzfc\textsuperscript{26}, supporting our interpretation.

Our model for the role of interfacial solvent reorganization in the hydrogen adsorption reaction bears some resemblance to a model for electrochemical proton
transfer suggested many years ago by Pecina and Schmickler\textsuperscript{27}. Using Monte Carlo simulations they showed that the activation energy for proton transfer increases at high interfacial fields due to the effect referred to above (at low field, the activation energy in their simulations in fact lowers due to a favourable configurational effect). More recently, Rossmeisl \textit{et al.}\textsuperscript{28} also considered models for the pH dependence of electrochemical proton transfer barriers, and emphasized the pH dependent contribution of the configurational entropy of the proton as the origin of the pH dependent kinetics. Our model emphasizes, on the other hand, the pH dependent enthalpy of interfacial water reorganization, as related to the pH dependent potential of zero charge of Pt on the RHE scale.
Figure 5. a) Cyclic voltammograms for Pt(111) in 0.1 M NaOH under argon atmosphere, registered at a scan rate of 50 mV.s$^{-1}$. The black line shows the blank of the bare electrode, whereas the blue line shows the Pt(111) electrode decorated with 0.17 ML of Ni(OH)$_2$, growth with instant deposition from a solution containing 5 mM Ni(NO$_3$)$_2$ (Ni$^{2+}$) and 3 cycles in the 0.1 M NaOH solution. b) Laser-induced coulostatic potential transients collected for the Pt(111) electrode. c) Laser-induced coulostatic potential transients collected for the Pt(111) electrode decorated with 0.17 ML of Ni(OH)$_2$. Dotted lines mark zero mV. The cartoons illustrate the average orientation of water molecules in the interface.
4.4. Conclusions

Given the experimental evidence presented in this paper, we conclude that a pH-dependent shift in the H-binding energy does not explain the significant pH dependence of the HER overpotential on Pt(111). We have studied its kinetic origin by a thorough interpretation of combined impedance measurements, Tafel plot analysis and laser-induced temperature jump experiments of interfacial solvent reorganization. We showed that not only the HER is slower in alkaline media, but also the H-UPD adsorption is slower, even if the thermodynamic driving force is the same for the different pH values on the RHE scale. Also the rate-determining step in the mechanism for the hydrogen evolution reaction is affected by the pH, in agreement with the observation that hydrogen adsorption becomes a kinetically hindered step in alkaline media.

We explain our observations by introducing a new model for the HER highlighting the role of interfacial water reorganization in the hydrogen adsorption step. This model is based on the experimental result that in acidic media, the H-UPD and HER occur close to the potential of zero charge, whereas in alkaline media, the H-UPD and the hydrogen evolution occur far from the potential of zero (total) charge. This would imply that in alkaline media the interfacial water network at the potential of H-UPD and HER interacts strongly with the strong interfacial electric field and is therefore more rigid and more difficult to reorganize during the charge transfer through the electrical double layer. This causes the hydrogen step to be slow in alkaline media. The presence of Ni(OH)$_2$ in submonolayer amounts improves the H-UPD charge transfer and the hydrogen evolution kinetics in alkaline pH by inducing a shift in the potential of total zero charge toward the HER equilibrium potential, as confirmed by laser-induced temperature jump experiments. In this new model, Ni(OH)$_2$ promotes the hydrogen evolution reaction by lowering the energy barrier necessary for the reorganization of the interfacial water network, allowing a more efficient proton/hydroxide transfer through the double layer. Importantly, the new model also implies that the hydrogen binding
energy is not the sole descriptor for the HER kinetics. The strength of the interfacial electric field, as determined by the distance of the potential of zero (total) charge (or the potential of maximum entropy) from the equilibrium potential of the reaction to be catalysed, impacts on the rate of hydrogen adsorption by influencing the energetic barrier associated with the reorganisation of interfacial water during charge transfer.

4.5. Acknowledgments

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4.6. References


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