1. Introduction

During the past couple of decades, electrochemistry received significant attention due to its relevance for devices and systems utilized for renewable energy conversion and storage.\(^1\)–\(^4\) Special attention was dedicated to the development of various electrode materials for electrolyzers,\(^5\) rechargeable batteries,\(^6\) fuel cells,\(^7\) etc., as well as to the fundamental understanding of processes at electrode/electrolyte interfaces.\(^8\)–\(^12\) A particularly important research area, that creates links between the fundamental understanding of materials’ properties and their functionality as electrode materials for energy conversion, is (electro)catalysis.\(^13\)

The comprehension of the nature and/or of the driving principles behind (electro)catalytic reactions has one of the prevailing roles in designing advanced interfaces for electrochemical energy conversion and storage, which suggests that electrocatalysis has simultaneously a remarkable scientific as well as economic importance.\(^14\)

A conceptual shaping of the theory of electrocatalysis was predominantly driven by the point of view formulated by Trasatti: “A true theory of electrocatalysis will not be available until activity can be calculated \textit{a priori} from some known properties of the materials”.\(^15\) Accordingly it was expected that some bulk or surface property of the material could be a descriptor of the (electro)catalytic activity.\(^16\)–\(^18\) The resulting models were usually “volcano”-type relations: (a) the Gschneidner-type of “volcano”-type plots which are illustrating changes in some physico-chemical property (e.g. cohesion energy) as a function of electronic structure or position of element in periodic system and (b) the Balandine-type of volcano-plots which are illustrating the Sabatier principle, \textit{i.e.} the “not too strong not too weak” binding of intermediates as a criterion for high reaction rate.\(^19\) Consequently, numerous catalytic descriptors were proposed over the years.
Practically, all introduced descriptors were related to the electronic structure of the electrode material. Some of the descriptors even delineated links between bulk and surface properties.20–22

However, interactions between the electrode material and the electrolyte could alter the electrode surface in a sense that no individual property of the material could be directly linked to the reaction rate. Consequently, this suggests not only that the electrolyte plays a very important role in the kinetics of electrocatalytic reactions,20,22 but rather emphasizes the importance of the interface itself and of the interfacial properties, which are strictly the result of the interaction between electrode material and electrolyte.23 In this study we investigate the hydrogen evolution reaction (HER) as model reaction, due to historical and practical significance for electrochemistry and due to its ‘simplicity’. Additionally, a conviction is that an in-depth understanding of HER as two electron process with only hydrogen evolution reaction (HER) as model reaction, due to historical and practical significance for electrochemistry and due to its ‘simplicity’. Additionally, a conviction is that an in-depth understanding of more complex reactions (e.g. four electron processes like oxygen reduction or oxygen evolution which proceed through at least two intermediates).

An important unresolved issue of contemporary interest related to the electrocatalysis of HER is the character of the relation between the exchange current (reaction rate at equilibrium potential) for HER and the M–H\text{ad} bond strength. This relation was usually perceived as “volcano”-type,17,24–27 where for the \( \Delta G(\text{M–H}{}_{\text{ad}}) \approx 0 \) (“not too strong not too weak” binding) the reaction rate will have its maximal value. Alternatively, in rare cases this relation was understood as linear, where the reaction rate increases the more negative the adsorption energy of the proton is.28,29 None of this two fundamentally different and confronting points of view (“volcano”-plot vs. linear-plot) has unambiguous experimental validation. The latter would imply that the Sabatier principle is not the only driving principle in electrocatalysis, contrary to gas-phase heterogeneous catalysis. Beyond the scientific debate “volcano” vs. linear, the knowledge about the true dependence between the reaction rate and the adsorption energy of the intermediate has also significant practical relevance. Namely, a linear dependence opens a perspective to discover/design catalysts with a superior activity to the previously perceived as the “top-of-volcano” one. Importantly, theoretical studies on this topic are not complementary to each other,17,28 while some rare experimental studies are in fact contradictory.26,29

In this work we tackle the essential difficulty in resolving the “volcano” vs. linear dependence controversy. Namely we devise an experimental procedure to measure an interfacial parameter that can be related directly to the adsorption energies of intermediates at the electrode/electrolyte interface. For this reason, we propose derived interfacial parameter as a new activity descriptor for HER. The practical and scientific value of the introduced interfacial descriptor is that it can be easily evaluated experimentally and can be also placed in an adequate theoretical framework, what is essential for the understanding of the activation process of HER. Importantly, this new descriptor allows the establishment of a functional relationship between properties of the material, properties of the interface and the reaction rate.

2. Experimental part

A commercial Scanning-Kelvin-Probe (SKP, K&M Soft-Control) utilized with an atmosphere and humidity control unit was used to obtain the work function of a group of polycrystalline samples (Au, Pd, Pt, Cu, Rh, Ir, Ru). Details of the experimental setup and the operational principles of SKP were given previously.30 Briefly, the samples were polished using a SiC paper (4000 grit), washed with de-ionized water and immediately dried under a stream of argon. Subsequently, the samples were transferred inside the measurement chamber, which is initially under nitrogen atmosphere and low relative humidity (1% RH, referred in the following as dry condition). After achieving steady atmospheric and humidity conditions the measurement was started. An array of 500 µm × 500 µm was measured, giving an array of 100 measurement points per sample, and the change in work function was measured continuously. The measurement under hydrogen atmosphere was performed by using 5% hydrogen in nitrogen, and to obtain the effect of humidity on the work function a maximum humidity of 95–98% was set (referred in the results as humid). Prior to the measurement the SKP tip was calibrated using a Cu/CuSO\text{4} (sat.) electrochemical couple.

3. Results and discussion

3.1. Which processes activate proton reduction?

The reaction pathway of HER is initiated with discharge of the proton known as Volmer step (reaction (1)), whose exact mechanism still remains a fundamental challenge. The proton discharge is followed either by the Heyrovsky step (reaction (2a)) or the Tafel step (reaction (2b)), both being recombination steps, where the former requires only one active site while the latter requires two active sites to produce one molecule of hydrogen.

\[
\begin{align*}
H^+ + e^- & \rightleftharpoons \text{H}_\text{ad} \quad \text{Volmer step} \quad (R1) \\
\text{H}_\text{ad} + H^+ + e^- & \rightleftharpoons \text{H}_2 \quad \text{Heyrovsky step} \quad (R2a) \\
2\text{H}_\text{ad} & \rightleftharpoons \text{H}_2 \quad \text{Tafel step} \quad (R2b)
\end{align*}
\]

The usual point of view is that the reaction rate is predominantly influenced by the adsorption energy of the intermediate H\text{ad}.17,25 However, the adsorption energy of H\text{ad} is not the only contributor to the activation barrier. In that context, it is essential to comprehend what really contributes to the free energy of activation (\( \Delta G^* \)) during the proton reduction. Consequently, it is essential to postulate what quantities define the transition state M·e····S·H\text{2}O\text{5−}, where M stands for metal surface; e\text{−} for electron, S for solvent and H\text{2}O\text{5−} for hydronium ion. The transition state is influenced by four almost simultaneous processes31 and their corresponding quantities: (a) the free energy of desorption of the water/solvent from the metal surface (\( \Delta G_{\text{desolv}} \)) as a precondition for the proton approach to the metal surface and for the electron transfer from the metal to the proton, (b) the work to transfer the solvated proton from the outer to the inner Helmholtz plane (W\text{m})), (c) the free energy of partial desolvation of the proton (\( \Delta G_{\text{desolv}} \)) or alternatively the energy required to deform...
respectively. The free energy of water desorption from the metal
surface, but rather prerequisite for the adsorption. This term evidently
depends on the nature of the metal. Different metals will have
different potential drops in the double layer (Galvani potential
difference), as well as different structuring of interfacial water
dipoles. However, considering that the dielectric constant in
the double layer for most of the metals is more less the same,
it is reasonable to assume that the resistance to transfer the
proton from the OHP to the IHP including its partial desolvation
will be approximately the same for different metals.

Being aware of the importance of the interfacial reorganization
of the solvent, it is worth to keep in mind that the proton
combines with a non-bonding electron pair of the water molecule
forming $\text{H}_2\text{O}^+$ with an enthalpy of formation of $-754$ kJ mol$^{-1}$.
The $\text{H}_2\text{O}^+$ cation attracts three additional $\text{H}_2\text{O}$ molecules in the
hydration shell and forms a time-averaged $\text{H}_3\text{O}^+$ cation with an
additional stabilization enthalpy of $-356$ kJ mol$^{-1}$. The latter
value also includes the rigid and diffuse hydration shells that
surround $\text{H}^+$. Clearly, even the partial desolvation of the proton
has to be an energetically very demanding process. This leaves an
open question of how exactly is HER triggered. In that context an
especially interesting fact is that the experimentally derived
activation energies have the same order of magnitude as non-
covalent bonds (hydrogen bonds etc.), which suggests that the
activation energy is strongly related to the rearrangement of
noncovalent bonds around the proton during the approach of
the proton to the electrode surface. Furthermore, this implies the
general importance of the solvent for the adsorption process in
inner sphere electrocatalytic reactions. This fact is usually
neglected in the literature. Just as a reminder, the reorganization
energy of the solvent is emphasized to be the key parameter that
determines the activation energy of the outer sphere redox
reactions as postulated by the Marcus theory of electron transfer.$^{32}$
Partial desolvation of the proton (including fluctuations and
deposition in the solvation sphere) is not exactly the same as the
classical reorganization energy of the solvent, although a
parallel can be drawn. Importantly, the group of Schmickler made
some estimations of the solvent coordinate during the proton
discharge,$^{33}$ while very recently Koper et al.$^{34}$ stressed the
importance of interfacial water reorganization for HER.

The fourth term in eqn (1), namely the energy to transfer the
proton from the Fermi level of the metal to the OHP of the
double layer ($\Phi^*$) through the metal/solvent interface, can be
directly related to the work function of the metal in vacuum
(check ESIf$^1$). From here we can conclude that the activation
energy for the HER is increasing with an increase of the work
function, which should be intuitive considering the definition
of the work function as the minimum work required for
extracting an electron from the Fermi level of the metal to
a position just outside the metal (far enough to eliminate
contributions from image forces). However, the experimental
linear relation between the work function of the metal and the
exchange current for HER, given previously by Trasatti$^{26}$
indicates the opposite trend. Namely, the increase in the work
additional energy. This suggests that the deformation of the
solvation sphere or in other words the interfacial reorganization
of the proton solvation shell is actually dictating the overall
value of the $(W_{\text{H}^+} + \Delta G_{\text{desolv}})$ term. This term in principle also
depends on the nature of the metal. Each of these parameters is complex and difficult for experi-
mental estimation or theoretical prediction, however, we can
discuss eqn (1) on a qualitative level. A graphical illustration of
the concept given by eqn (1) is shown in Fig. 1.

While all four processes are contributing to the activation
barrier, only the sum of the last three terms $(W_{\text{H}^+} + \Delta G_{\text{desolv}} + \Phi^*)$
practically represents the free energy of adsorption of the proton.
They can be understood as the interaction of the proton with the
electric field in the double layer, the interaction of the proton
with the solvent and the interaction of the proton with the metal,
respectively. The free energy of water desorption from the metal
surface, $\Delta G_{\text{desolv}}$, is not integral part of the adsorption energy,
but rather prerequisite for the adsorption. This term evidently
strongly depends on the nature of the metal.

The work to transport the solvated proton from the outer to
the inner Helmholtz plane $(W_{\text{H}^+})$ and the free energy of partial
desolvation of the proton $(\Delta G_{\text{desolv}})$ are mutually dependent.
Namely, if the deformation of the solvation sphere is an easy
going process then the work required to transport protons from
OHP to IHP will be relatively low. However, if the deformation
of the solvation sphere is energetically demanding then the
proton shift from the OHP towards the IHP will require

\[
\Delta G^* = \Delta G_{\text{desolv}} + \{(W_{\text{H}^+} + \Delta G_{\text{desolv}}) + \Phi^*\}
\]

(1)

Fig. 1 Graphical illustration of the processes (explanation given in the
text) in the double layer necessary for the reduction of the proton. Red
circles denote hydrogen atoms, blue circles represent oxygen atoms and
smaller red circles surrounded each by four water molecules represent
solvated proton $(\text{H}_2\text{O}_4^+)$. 

The proton shift from the OHP towards the IHP will require

\[
\Delta G^* = \Delta G_{\text{desolv}} + \{(W_{\text{H}^+} + \Delta G_{\text{desolv}}) + \Phi^*\}
\]

(1)
function of the metal was related to higher exchange currents for the HER. The observed experimental relation between the exchange current for HER and the work function of metals was understood as a “secondary effect”. It was observed that a high value of the work function of metal corresponds to more positive values of the $E_{pex}$ (linear relation with slope around 0.8). In this way, the high value of the work function and/or positive values of $E_{pex}$ assure high concentration of protons in the double layer at the reversible potential for hydrogen evolution. Consequently, the Volmer step ($\text{H}^+ + e^- = \text{H}_ad$) which initiates the HER should be very facile, requiring almost no overpotential. This indeed can be the case, if the electron transfer rate for all studied metals is substantially higher than the proton transfer rate. In other words, if the electron transfer for a given metal is more sluggish than the electron transfer rate for some other metal with lower work function, this will have no effect on the proton reduction rate, because the reaction is limited by the proton transfer rate. Formally, from the point of the rate law this would mean that although an increase in the work function reduces the exponential factor (enlarges activation free energy) it enlarges the preexponential factor even more, so that the overall exchange current increases with an increase of the work function of the metal.

3.2. Changes in the properties of the double layer in relation to the preexponential factor of the rate law

If one compares the kinetics of HER on Pt to the kinetics of outer sphere reactions (e.g., $\text{Fe}^{3+}/\text{Fe}^{2+}$ redox reaction) on some metal, it is noticeable that despite the very similar activation enthalpies, the exchange current for HER is significantly lower. This suggests that for inner sphere electrocatalytic reactions the preexponential (frequency) factor is much lower.

While the drift speed of protons in the bulk of electrolyte is $36.23 \times 10^{-6}$ m s$^{-1}$, in the electric double layer (electric field is in the range $10^6$–$10^7$ V m$^{-1}$) it reaches $3.623$–$36.23$ m s$^{-1}$ (mobility of proton in water at $T = 298$ K is $36.23 \times 10^{-8}$ m$^2$ V$^{-1}$ s$^{-1}$, in case of a typical potential difference of 1.0 V applied between two electrodes separated by a distance of $d = 10^{-2}$ m, where the electric field is $10^7$ V m$^{-1}$). If one considers that the dielectric constant of the water molecules in the double layer is additionally reduced by one order of magnitude in comparison to the bulk, the drift speed of protons should be even higher, what should enlarge the number of attempts on the activation barrier per unit of time. This is usually denoted in the preexponential factor of the rate law. The usual expression for the rate constant ($k$) is:

$$k = A \exp \left( \frac{-\Delta G^*}{RT} \right) \quad (2)$$

where $A$ is the preexponential frequency factor, $T$ is the temperature, $R$ the universal gas constant and $\Delta G^*$ is the activation free-energy. In the case of outer sphere reactions where reactants and products stay at OHP, the preexponential factor formally does not depend significantly on the nature of the electrode material. However, during the HER the electric field shifts the reactant (proton) from the OHP to the IHP of the double layer and only protons which reach the IHP will react. Evidently, the approach of the proton to the surface strongly depends on the structure of the double layer which again depends on the arrangement of the interfacial water dipoles. The arrangement of interfacial water dipoles originates in the metal–water interactions, meaning that the preexponential factor of the inner sphere electrocatalytic reactions including HER, depends on the nature of the electrode material. If the drift speed of the proton is $36.2$ m s$^{-1}$ and the proton is shifted from the OHP towards the IHP over a distance of around $0.1$ nm (approximately half of the thickness of the double layer) the frequency of attempts of protons on the activation barrier is around $3.6 \times 10^{11}$ Hz.

A semi-classical treatment of electron transfer\textsuperscript{39,40} suggests that the electron transfer rate is:

$$k = k_{el} \Gamma_n \nu_n \exp \left( \frac{-\Delta G^*}{RT} \right) \quad (3)$$

where $k_{el}$ is the electronic transmission coefficient ($k_{el} \leq 1$) related to the probability of electron-tunnelling and often is taken to be unity for a reaction where the reactant is close to the electrode, so that there is strong coupling between the reactant and the electrode; $\Gamma_n$ is the nuclear tunneling factor ($\Gamma_n \geq 1$) which corrects the electron-transfer rate for reactants (protons) that react without fully surmounting the classical free-energy barrier; $\nu_n$ is the nuclear frequency factor which represents the frequency of attempts on the energy barrier and is generally associated with bond vibrations and the solvent motion.

In relation to the above discussed acceleration of protons in the double layer by more than 6 orders of magnitude in comparison to the bulk of electrolyte, we expect that the larger the acceleration is, the larger will be the nuclear frequency factor or the frequency of attempts on the energy barrier. At the same time, the larger the acceleration is, the closer to the electrode surface will be the positioned proton, so the higher will be the tunnelling probability for both, electrons and protons at the applied electrode potential. Considering that metals with high work function have the double layer highly saturated with protons, it seems reasonable to assume that metals with high work function will have higher preexponential frequency factor, because the probability that an active site will interact with a proton is higher.

3.3. The adsorption through the water layer – the change in the work function as an in situ indicator of the metal–hydrogen bond strength

An important segment for understanding the HER catalytic trends is the interdependence between estimated values of exchange currents (kinetic parameter), work functions (property of material) and adsorption energies of $\text{H}_ad$ intermediates (interfacial parameter). While the values of the work function of the metals and the corresponding exchange currents for HER are relatively reliable,\textsuperscript{26} it seems that previous evaluations of the adsorption energies of $\text{H}_ad$ intermediate contain certain inconsistency. In principle, the high work function should be linked with weakening of the M–H bond (more positive adsorption energy of $\text{H}_ad$ intermediate), according to eqn (1). In fact, some works indicate that (in the case of d-metals) this kind of
reasoning is justified, although the estimated values of adsorption energies as well as the given trends seem to be inconsistent or even contradictory. If we recall that the sum of the last three terms in eqn (1) \( \Delta G_{\text{desolv}} + \Delta \Phi^* \) is analogue to the free energy of adsorption of proton, and considering the complexity and the mutual interdependence between the terms in eqn (1) it seems that a straightforward evaluation of adsorption energies of the \( \text{H}_\text{ad} \) intermediate would be necessary and would require a reliable experimental approach.

An experimental identification and/or confirmation of an appropriate activity descriptor is usually the missing link between theoretical concepts in electrocatalysis and experimentally derived activity trends. In that context, as stated above, the impact of the work function on the electrocatalysis of HER is more than just a "secondary effect" (or double layer effect). We propose that the work function has major implications on the proton and the electron transfer rate not only via the preexponential frequency factor but also through the exponential activation factor, which is strongly influencing the M–H bond formation. From eqn (1) the adsorption energy \( \Delta G_{\text{ad}} = \Delta W + \Delta \Phi \) becomes more endergonic (weaker bonding) when the work function is increasing. This view can be supported by the work of Conway and Bockris who gave a quantitative link between the adsorption energy (or bond strength) and the work function (as a measure of the cohesion energy in the metal) illustrated with the Pauling equation:

\[
E(\text{M–H}) = 0.5[D(\text{M–M}) + D(\text{H–H})] + 23.06(X_{\text{M}} - X_{\text{H}})^2
\]

where, \( E(\text{M–H}) \) – the metal–hydrogen bond strength, \( D(\text{M–M}) \) – the dissociation energy of the M–M bond in the metal lattice, \( D(\text{H–H}) \) – the dissociation energy of the hydrogen molecule, \( X_{\text{M}} \) – the electronegativity of the metal and \( X_{\text{H}} \) – the electronegativity of hydrogen. In Fig. 2 is shown the possible relationship between the work function (of d-metals) and the M–H bond strength. From the tendency given in Fig. 2 one can conclude that stronger adsorption of \( \text{H}_\text{ad} \) could result in more pronounced drop in work function.

A specific feature of the electrochemical environment is that the adsorption proceeds through the interfacial water layer. Although some authors assumed significant changes in the work function when comparing vacuum conditions with conditions of metal covered by water, reports on these phenomena are contradictory. Elsewhere, work functions of metals in a solution were considered practically to be the same as those of metals in a vacuum. It was assumed that the water behaved like a dielectric continuum that shifted the absolute scale of reactivity to the same extent for all metals due to water dipoles, which alter the surface potential at the solid/liquid interface. The first step in our experimental sequence suggests absence of significant change in the work function under humid conditions, employing in situ experiments based on SKP (scanning Kelvin probe). The work function in dry nitrogen atmosphere changed only slightly compared to humid nitrogen atmosphere. In Fig. 3 after introducing humidity for most of the metals a slight drop in the work function is noticeable except for Cu and Ru, where a slight increase in the work function is observed. This can be a result of the fact that initially it is very difficult to remove water completely from the metal surface. However, surprisingly the measured values of the work function match to the literature reported ones for the vacuum conditions. This can be a consequence of the fact that interfacial water layer has extremely low dielectric constant (\( \varepsilon \approx 4 \)), almost 20 times lower than liquid water, forming a structure known as "ice 7", without a preferential water dipole orientation. This can be additionally supported by works of Thiel et al. where it was shown that the impact of the interfacial water layer on the change in the work function is not so straightforward and remains very difficult to predict. It appears that, in contrast to what is usually believed, at open circuit conditions the associative (molecular) adsorption of water through oxygen does not cause a significant drop in the work function. For example for Pt immersed in aqueous electrolyte the open circuit
potential is around 0.9 V$_{\text{RHE}}$ which (according to eqn (5)) responds to 5.35 eV on absolute scale.

$$E(\text{SHE})/V = \frac{\Phi}{e} - 4.44$$  \hspace{1cm} (5)

In fact, the associatively adsorbed water often exhibits a variety of adsorption geometries to accommodate the intermolecular hydrogen bonding. Some of these geometries may have also oxygen atoms pointing away from the surface. At the same time, at metal surfaces where water is dissociated to form OH (or even O), even an increase in the work function is expected. Finally, depending on the electrode potential, water dissociation may also generate atomic hydrogen whose presence at the metal surface can cause both, drop or increase in the work function. 48

In the next experimental step, the introduction of hydrogen gas (through water adsorbed at the surface) caused a drop in the work function for all analysed metals, except Cu. The drop in the work function spans from 0.0 eV in the case of Cu till 0.5 eV in the case of Ru and can be due to the dissociation of molecular hydrogen on the metallic surface. This is in line with the earlier established linear relation between change in the work function and adsorption energy of atomic hydrogen, introduced by Jaksic et al. 19

$$\log j_0 \propto \frac{\beta \Delta G_{\text{ad}}}{k_B T} \propto \frac{\beta r \Delta \Phi}{k_B T}$$  \hspace{1cm} (6)

where $j_0$ – exchange current, $\beta$ – symmetry factor of activation barrier, $\Delta G_{\text{ad}}$ – change in the adsorption energy of intermediate in the rate determining step, $\Delta \Phi$ – change in the work function upon adsorption of intermediate, $r$ – the slope of the apparent experimental linear dependence between $\Delta G_{\text{ad}}$–$\Delta \Phi$ (other quantities were defined previously). From this relation stems that the stronger the hydrogen adsorption is the larger the drop in the work function. Interestingly, for Cu the change in the work function was negligible, which would imply that $\Delta G_{\text{ad}} \approx 0$ (thermoneutral adsorption). This, according to some authors, should be criteria of superior activity. 17,24,25,37 However, Cu is out of this seven analyzed metals by far the worst electrocatalyst for HER. Metals with optimal adsorption energy of hydrogen will have the highest equilibrium concentration of protons in the thin water layer. Those metals who bind hydrogen too weakly will have very low surface concentration of atomic hydrogen and therefore also a low concentration of solvated protons, while those who bond hydrogen strongly will have a high surface concentration of hydrogen, but no possibility to desorb hydrogen into the water layer in a form of protons. The values of the measured work function can be related to the electrode potential on the standard hydrogen scale (eqn (5)). 79

From eqn (5) stems that, for example Pt that exhibits the highest work function upon adsorption of hydrogen through the water layer, will have the most positive potential on standard hydrogen scale. This is an indicator of acidification of the thin water layer and a consequence of the optimal binding energy of hydrogen.

In the last experimental step, the hydrogen atmosphere humidity was reduced to zero, which caused a partial recovery of the work function for Ru, Ir, Pd and Rh. In the case of Cu, Au and Pt, removal of the humidity did not induce significant change in the work function in comparison to the previous state. A complete recovery of the initial values of work functions would require substantial time.

Interestingly, it was previously shown by Trasatti that the adsorption enthalpy of hydrogen exhibits opposite trends when comparing adsorption from the gas-phase with adsorption from the solution. 30 In the same work those metals which strongly adsorb hydrogen from the gas-phase weakly absorb hydrogen from the solution. From the tendencies shown in the Fig. 3 it seems that although water does not change the work function of the analysed metals significantly, it behaves as a promoter of hydrogen adsorption for some metal surfaces. This phenomena could not be explained in this instance. Nevertheless, underlying reasons could be the impact of water dipoles and the electric field on the adsorption 31 by effecting metal–water interactions 52 and noncovalent bonding in the hydrogen-bonding network. 53 Consequently, the presence of water dipoles at the metal surface and the existence of an electric field in the double layer, dependently from the properties of “ice”-like structure in the double layer 46,47 can promote or inhibit the metal–hydrogen bond formation. From here, it becomes evident that one of the central tasks in future experimental and theoretical studies related to electrochemical interfaces is to understand the role of the interfacial water molecules/dipoles in reaction mechanisms and its impact on electrode reaction rates. This understanding would be particularly important for reactions that are perceived to be inner sphere electrocatalytic reactions. 54

Further, the significance of our approach also becomes evident when one tries to establish structure–activity relations, using the bulk or surface properties of materials. If one considers the recent works of Koper et al., 18 it has been shown that, for the transition metals, the number of outer electrons is in good relation with the adsorption energies. From Fig. 4 it can be observed that the most active metal for the HER (Pt) has a number of outer electrons equal to 10. At the same time from Fig. 4 it can be observed that the change in the work function upon adsorption of hydrogen through the water layer for the most active metals has the optimal value of around 0.35 eV. Additionally, from Fig. 4 one can also notice that for the same number of outer electrons (electronic structure parameter) the drop in the work function can be significantly different (example of Au and Cu), which confirms that the individual bulk or surface properties can be problematic as catalytic descriptor, as is previously stated.

Utilizing the change in the work function upon adsorption of hydrogen through the water layer as the interfacial activity descriptor we obtain the “volcano”-plot shown in Fig. 5. The top of this experimental “volcano” indicates the optimal interaction of the metal with hydrogen in the presence of water.

Evidently, we can use the change in the work function upon adsorption of hydrogen through a water layer as catalytic descriptor.
energies of these four metals were very similar to each other. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor. In fact, the adsorption energies were used as descriptor.

The experimental derivation of $\Delta \Phi$ is straightforward and can be conveniently used also for the validation of values and trends of computationally derived adsorption energies in electrolyte at open circuit conditions.

From Fig. 5 we can observe that the introduced descriptor is much more sensitive than the previously estimated adsorption energies. For example the values of the exchange currents for Pd, Rh, Ir and Ru (Fig. 5) are clearly linearly scaled with the introduced descriptor which was not the case when adsorption energies were used as descriptor. In fact, the adsorption energies of these four metals were very similar to each other although their exchange currents could differ for more than one order of magnitude. From Fig. 5 we can notice that the bonding of hydrogen on Pt is slightly weaker than the optimal which is opposite to previous reports. The slope of the ascending branch of the “volcano” shown in Fig. 5 is around 122 mV dec$^{-1}$ which is very close to 118 mV dec$^{-1}$, the theoretical Tafel’s slope of HER when the discharge (Volmer) reaction is the rate determining step (weakly binding metals). The descending branch of the “volcano” has similar value to the slope that responds to the value of the theoretical Tafel’s slope of HER when the recombination step is the rate determining for the case when the coverage with the intermediate is close to unity (strongly binding metals).

Summing up the most important experimental and theoretical findings we can state the following:

1. The demonstrated experimental “volcano”-plot resembles the “volcano” curves reported previously where the optimal value of adsorption energy allows the highest exchange current density. Our experimental “volcano”-curve is an excellent experimental confirmation of the Sabatier principle in electrocatalysis. However, the descriptor used in Fig. 5, analogue to the adsorption energy, is based on the last three terms of eqn (1). Importantly, the first term that denotes the free energy of desorption of water/solvent from the metal surface ($\Delta G_{\text{desorp}}$) as precondition for proton approach to the metal surface and electron transfer from metal to proton, is not part of the descriptor although it contributes to the activation energy.

2. The higher the work function for an electron–donor cathodic reaction such as HER, which proceeds at potentials more negative than potential of zero charge, the lower the electron transfer rate should be. However, the electron transfer step is not the rate limiting process for HER. It seems that overcoming of the proton transfer barrier is one of the central processes that determine the exchange current for HER. In this context of particular importance, it will be challenging in the future to address phenomena of proton transfer, especially possible conditions of proton tunneling.

3. The purpose of this work is to bring a conceptual novelty which will contribute to a better understanding of HER. Therefore we focused initially only on noble metals, materials whose work functions and exchange currents are straightforward to obtain. However, future work would proceed with applying the presented approach for the analysis of HER on various single crystal facets, defected facets and nanostructures as well as on investigating materials which are considered to be realistic alternative to Pt (various alloys, chalcogenides, phosphides etc.). Finally, this approach can be extended to other important reactions like oxygen reduction reaction etc.

4. Conclusion

In this work it is demonstrated that the experimentally determined change in the work function upon adsorption of hydrogen through a water layer is a realistic indicator of metal–hydrogen interaction during the electrocatalytic proton reduction. The interfacial descriptor introduced can be used as a tool to evaluate theoretically established values and trends in the adsorption energies. Additionally, this descriptor could be utilized as a tool...
to validate models proposed by the computational chemists. In more general terms, it was shown that interfacial parameters, that include the metal–solvent–proton interaction, are the most relevant/realistic descriptors of the electrocatalytic activity because they take into consideration the interaction between the electrode and the electrolyte, something that cannot be easily predicted using computational methods.

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References

1 I. Katsounaros, S. Cherevko, A. R. Zeradjanin and K. J. J. Mayrhofer, Angew. Chem., Int. Ed., 2014, 53, 102–121.
2 N. S. Lewis and D. G. Nocera, Proc. Natl. Acad. Sci. U. S. A., 2006, 103, 15729–15735.
3 J. O. Bockris, Science, 1972, 176, 1323.
4 J. O. M. Bockris, Int. J. Hydrogen Energy, 2013, 38, 2579–2588.
5 J. Suntivich, K. J. May, H. A. Gasteiger, J. B. Goodenough and J. O. M. Bockris, Acc. Chem. Res., 2013, 46, 1053–1061.
6 V. R. Stamenkovic, B. Fowler, B. S. Mun, G. Wang, P. N. Ross, C. A. Lucas and N. M. Markovic, Science, 2007, 315, 493–497.
7 J. O. Bockris, J. Electrochem. Soc., 1984, 131, 290.
8 J. Matsumoto and E. Sato, Mater. Chem. Phys., 1986, 14, 397–426.
9 H. Dau, C. Limberg, T. Reier, M. Risch, S. Roggan and P. Strasser, ChemCatChem, 2010, 2, 724–761.
10 D. Strmcnik, M. Uchimura, C. Wang, R. Subbaraman, N. Danilovic, D. van der Vliet, A. P. Paulikas, V. R. Stamenkovic and N. M. Markovic, Nat. Chem., 2013, 5, 300–306.
11 A. R. Zeradjanin, A. A. Topalov, Q. Van Overmeere, S. Cherevko, X. Chen, E. Ventosa, W. Schuhmann and K. J. J. Mayrhofer, RSC Adv., 2014, 4, 9579.
12 J. Greeley and N. M. Markovic, Energy Environ. Sci., 2012, 5, 9246.
13 A. R. Zeradjanin, E. Ventosa, A. S. Bondarenko and W. Schuhmann, ChemSusChem, 2012, 5, 1905–1911.
14 J. X. Wang, F. A. Uribe, T. E. Springer, J. Zhang and R. R. Adzic, Faraday Discuss., 2009, 140, 347–362.
15 J. K. Nørskov, J. Rossmeisl, A. Logadottir, L. Lindqvist, J. R. Kitchin, T. Bligaard and H. Jónsson, J. Phys. Chem. B, 2004, 108, 17886–17892.
16 J. K. Nørskov, T. Bligaard, A. Logadottir, J. R. Kitchin, J. G. Chen, S. Pandelov and U. Stimming, J. Electrochem. Soc., 2005, 152, J23.
17 F. Calle-Vallejo, N. G. Inoglu, H.-Y. Su, J. I. Martínez, I. C. Man, M. T. M. Koper, J. R. Kitchin and J. Rossmeisl, Chem. Sci., 2013, 4, 1245.
18 J. M. Jakšic, N. M. Ristic, N. V. Krstajic and M. M. Jakšic, Int. J. Hydrogen Energy, 1998, 23, 1121–1156.
19 O. A. Petrii and G. A. Tsirlina, Electrochim. Acta, 1994, 39, 1739–1747.
21 P. Searson, P. Nagarkar and R. Laianision, Int. J. Hydrogen Energy, 1989, 14, 131–136.
22 A. R. Zeradjanin, A. A. Topalov, S. Cherevko and G. P. Keeley, Int. J. Hydrogen Energy, 2014, 39, 16275–16281.
23 A. R. Zeradjanin, N. Menzel, P. Strasser and W. Schuhmann, ChemSusChem, 2012, 5, 1897–1904.
24 H. Gerischer, Bull. Soc. Chim. Belg., 2010, 67, 506–527.
25 R. Parsons, Trans. Faraday Soc., 1958, 54, 1053.
26 S. Trasatti, J. Electroanal. Chem. Interfacial Electrochem., 1972, 39, 163–184.
27 M. T. M. Koper, J. Solid State Electrochem., 2016, 20, 895–899.
28 E. Santos, P. Quaino and W. Schmickler, Phys. Chem. Chem. Phys., 2012, 14, 11224.
29 S. Trasatti, in Proceedings of the Symposium on Electro-catalysis, ed. W. E. O’Grady, P. N. Ross and F. G. Will, The Electrochemical Society Proceedings Series, Pennington, N. J., 1982, p. 73.
30 M. Rohwerder and F. Turcu, Electrochim. Acta, 2007, 53, 290–299.
31 S. Harinipriya and M. V. Sangaranarayanan, J. Phys. Chem. B, 2002, 106, 8681–8688.
32 R. A. Marcus and N. Sutin, Biochim. Biophys. Acta, Rev. Bioenerg., 1985, 811, 265–322.
33 A. R. Zeradjanin, A. A. Topalov, S. Cherevko and G. P. Keeley, Phys. Rev. B: Condens. Matter Mater. Phys., 2009, 79, 235436.
34 I. Ledezma-Yanez, W. D. Z. Wallace, P. Sebastián-Pascual, V. Climent, J. M. Feliu and M. T. M. Koper, Nat. Energy, 2017, 2, 17031.
35 S. Trasatti, J. Electroanal. Chem. Interfacial Electrochem., 1971, 33, 351–378.
36 W. Sheng, H. A. Gasteiger and Y. Shao-Horn, J. Electrochem. Soc., 2010, 157, B1529.
37 D. Galizzioli and S. Trasatti, J. Electroanal. Chem. Interfacial Electrochem., 1973, 44, 367–388.
38 G. Jerkiewicz, Electrocatalysis, 2010, 1, 179–199.
39 M. J. Weaver and G. E. McManis, Acc. Chem. Res., 1990, 23, 294–300.
40 M. J. Weaver, Chem. Rev., 1992, 92, 463–480.
41 B. E. Conway and J. O. Bockris, J. Chem. Phys., 1957, 26, 532.
42 A. T. Kuhn, C. J. Mortimer, G. C. Bond and J. Lindley, J. Electroanal. Chem. Interfacial Electrochem., 1972, 34, 1–14.
43 P. Rüetschi, J. Chem. Phys., 1955, 23, 195.
44 S. Trasatti, Electrochim. Acta, 1991, 36, 1659–1667.
45 J. M. Heras and L. Viscido, Appl. Surf. Sci., 1980, 4, 238–241.
46 I. Danielewicz-Ferchmin, J. Phys. Chem., 1995, 99, 5658–5665.
47 M. F. Toney, J. N. Howard, J. Richer, G. L. Borges, J. G. Gordon, O. R. Melroy, D. G. Wiesler, D. Yee and L. B. Sorensen, Surf. Sci., 1995, 335, 326–332.
48 P. A. Thiel and T. E. Madey, Surf. Sci. Rep., 1987, 7, 211–385.
49 S. Trasatti, J. Electroanal. Chem. Interfacial Electrochem., 1986, 209, 417–428.
50 S. Trasatti, Russ. J. Electrochem., 2005, 41, 1255–1264.
51 S. A. Wasileski, M. T. M. Koper and M. J. Weaver, J. Am. Chem. Soc., 2002, 124, 2796–2805.
52 R. R. Nazmutdinov, M. Probst and K. Heinzinger, Chem. Phys. Lett., 1994, 222, 101–106.
53 D. Strmcnik, K. Kodama, D. van der Vliet, J. Greeley, V. R. Stamenkovic and N. M. Markovic, Nat. Chem., 2009, 1, 466–472.
54 J. Staszak-Jirkovský, R. Subbaraman, D. Strmcnik, K. L. Harrison, C. E. Diesendruck, R. Assary, O. Frank, L. Kobr, G. K. H. Wiberg, B. Genorio, J. G. Connell, P. P. Lopes, V. R. Stamenkovic, L. Curtiss, J. S. Moore, K. R. Zavadil and N. M. Markovic, ACS Catal., 2015, 5, 6600–6607.
55 J. Rossmeisl, K. Chan, E. Skúlason, M. E. Björketun and V. Tripkovic, Catal. Today, 2016, 262, 36–40.
56 R. R. Dogonadze and L. I. Krishtalik, Russ. Chem. Rev., 1975, 44, 938–945.
57 A. R. Zeradjanin, J.-P. Grote, G. Polymeros and K. J. J. Mayrhofer, Electroanalysis, 2016, 28, 2256–2269.
58 J. Tymoczko, F. Calle-Vallejo, W. Schuhmann and A. S. Bandarenka, Nat. Commun., 2016, 7, 10990.
59 J. D. Benck, T. R. Hellstern, J. Kibsgaard, P. Chakthranont and T. F. Jaramillo, ACS Catal., 2014, 4, 3957–3971.
60 Z. W. Seh, J. Kibsgaard, C. F. Dickens, I. Chorkendorff, J. K. Nørskov and T. F. Jaramillo, Science, 2017, 355, eaad4998.