Impact of additives on zirconium oxidation by water: mechanistic insights from first principles

Mikaela Lindgren* and Itai Panas

Zirconium alloys are widely used as cladding materials employed to contain the fission fuel in nuclear power plants. A limiting factor for fuel longevity is the corrosion property of the zirconium alloys. In the reactor, the main corrodent is water. The oxide forms thermodynamically during hydrogen evolution. Due to the corrosion mechanism, a fraction of the hydrogen is transferred to the alloy. It has long been known that the alloying elements actually control the hydrogen pick-up fraction, HPUF. A mechanism that explains these observations by means of density functional theory calculations is presented and validated. A hydroxylated grain boundary model decorated by various transition metal, TM, ions is employed to study the dependence of the hydrogen evolution reaction, HER, on the choice of TM ion and spin state along the hydride–proton recombination pathway. The efficiency of the system to utilize the overpotential for hydrogen evolution, originating from the overall corrosivity of the alloy, is found to be decisive for the HPUF. A dual origin of the detrimental effects of Co and Ni additives on the HPUF is identified.

I. Introduction

Today, fundamental aspects of apparently disparate branches of material science are seen to coalesce, which is due to a unifying atomistic paradigm in conjunction with the prevailing computational technology, as manifested in implementations of density functional theory. Here, this is realised in the analysis of corrosion, where electrochemical interpretive concepts have long since been employed. Thus, corrosion is associated with a redox process where the reduction is said to occur at the cathode, while oxidation takes place at the anode. Sustained corrosion requires media that conduct electrons and ions in order to maintain electroneutrality at both electrodes. Clearly, such a corrosion process is related to the operation of a fuel cell, yet the corresponding technological implications remain to be explored more fully.

The control of electron and ion transport is crucial both for obtaining improved materials for fuel cell applications and for improved corrosion protection; the same materials are often investigated for similar reasons. This is so for ZrO₂, which constitutes the protective barrier layer on zirconium alloys, e.g. the cladding for nuclear fuel. Indeed, the transport properties of this barrier oxide determine the durability of the fuel assembly itself. Zirconia based materials are also ubiquitous in fuel cell research owing to their properties as solid oxide proton conductors. The added value of such transferability between complementary disciplines is sought by clarifying and utilizing their common contextual framework. In this spirit, in a previous study, the oxidation of zirconium alloys by water was analysed. The process was subdivided into an anode process

\[ \text{Zr} + 2\text{O}^{2-} \rightarrow \text{ZrO}_2 + 4\text{e}^- \]  

(R1)

and a cathode process which eventually results in hydrogen evolution, occurring at transition metal sites in hydroxylated grain boundaries of ZrO₂ (see Fig. 1). Evidence for grain boundary hydroxides was provided by infrared spectroscopy, while transition metal decoration was reported by atom probe tomography. The cathode reaction was subdivided into two steps, the first of which is the formation of a three-centre hydride

Fig. 1 Representative structure for a model of hydroxylated grain boundary ZrO(OH)₂ in ZrO₂. Here, the grain boundary is decorated with Fe ions in oxidation state II. Oxygen is represented as red, zirconium as light blue, iron as purple and hydrogen as white.
\[ \text{TM}_{\text{GB}}^X + \text{Zr}^{IV} + \text{OH}^- + 2e^- \rightarrow \text{TM}_{\text{GB}}^X\text{H}^-\text{Zr}^{IV} + \text{O}^2^- \quad (\text{R}2\text{a}) \]

or the equivalent

\[ \text{TM}_{\text{GB}}^X + \text{Zr}^{IV} + \text{H}^+ + 2e^- \rightarrow \text{TM}_{\text{GB}}^X\text{H}^-\text{Zr}^{IV} \quad (\text{R}2\text{b}) \]

and the second is a subsequent hydrogen evolution reaction

\[ \text{TM}_{\text{GB}}^X\text{H}^-\text{Zr}^{IV} + \text{OH}^- \rightarrow \text{TM}_{\text{GB}}^X + \text{Zr}^{IV} + \text{H}_2 + \text{O}^2^- \quad (\text{R}3\text{a}) \]

or

\[ \text{TM}_{\text{GB}}^X\text{H}^-\text{Zr}^{IV} + \text{H}^+ \rightarrow \text{TM}_{\text{GB}}^X + \text{Zr}^{IV} + \text{H}_2 \quad (\text{R}3\text{b}) \]

where \( X = \text{II or III}, \text{and TM} = \text{Ti, V, Cr, Mn, Fe, Co, Ni, Cu or Zn}. \) The (R2a) and (R3a) emphasize that reactive protons reside on hydroxide ions. Support for the possible presence of hydride intermediates in hydroxylated grain boundaries was arrived at by comparing computed vibration frequencies to experimental infrared spectra. When II ions were present, the hydrides were found to form spontaneously for all elements investigated. In contrast, for TMIII the reaction (R2) becomes exothermic for Ti, V, and Cr, while it is endothermic for Mn, Fe, Co, and Ni (see Fig. 2).

The energy changes for the hydride–proton recombination reaction (R3) were computed. The resulting \( H_2 \) evolution energy versus choice of elements was plotted, i.e. for TiII, VIII, CrIII, MnII, FeII, CoII, NiII, CuII and ZnII, by using the findings for the oxidation state III (see Fig. 2) as a weight function.

For all 3d transition metals investigated, the 3d orbitals are the ones which interact directly with the hydride. To isolate the properties of pure zirconia, \( \text{Ca}^{2+} \) was introduced as a reference II ion replacing the 3d transition metal elements of the oxidation state II. Complementary to \( \text{Ca}^{2+} \), the impacts of inert ions, in oxidation states I and III, were also evaluated. This was achieved by replacing \( \text{Ca}^{2+} \) by \( \text{Na}^- \) and \( \text{Sc}^{3+} \), respectively. Thus, a stability test for the hydride–proton recombination energy of pure zirconia is provided, see shaded horizontal band in Fig. 3.

Finally, the correlation between the computed reaction energy changes for (R3) and the experimental ratios of hydrogen picked up by the alloy to the total amount of hydrogen converted, i.e. the hydrogen pick-up fraction, HPUF, was sought and striking agreement was obtained (see Fig. 3).

The purpose of the present study is to explore the cause of this agreement and its possible consequences. An analysis of the structural origins of the reaction energy changes in (R3) is included. Moreover, beyond the property of the reaction energy to offer a descriptor for predicting the HPUF, we attempt to unravel the reaction mechanism for the proposed hydride–proton recombination reaction. Finally, the origin and possible implication of the detailed correlation between the energy change associated with hydrogen evolution and the hydrogen pick-up fraction are discussed.

II. Modelling considerations

II.a. The model

The present study utilizes a periodically repeating hydroxylated grain boundary embedded in ZrO2. It was constructed by inserting one unit cell of monoclinic ZrO(OH)2 (ref. 6) \( (5.4 \text{ Å} \times 10 \text{ Å} \times 5.4 \text{ Å}) \) between two supercells of monoclinic ZrO2 \( (5.4 \text{ Å} \times 10.8 \text{ Å} \times 5.4 \text{ Å}) \), where the unit cell doubling is in the \( b \) direction. The two subcells were fused “epitaxially” by making contact between the \( \{a,c\} \) planes of each subcell. A periodically layered structure in the \( b \) direction was obtained. The starting ZrO(OH)2 structure was obtained by replacing Li+ with H+ in Li2ZrO3. Upon replacing Li+ by H+ and subjecting the system to geometry optimization, protons bind to O2− ions thereby forming hydroxide ions, hence the ZrO(OH)2 structure was formed. Interestingly, an excellent proton conductor has been obtained experimentally by performing ion exchange on Li2ZrO3, replacing Li+ with H+.

Although the stability of the model was extensively investigated, only a subset of all calculations for the system is provided here. The choice of grain boundary model is far from unique. However, for the model to offer a valid chemical representation of the hydroxylated interface between two ZrO2 grains, there are certain requirements that need to be satisfied.

Essential to the modelling, besides reproducing the large band gap signature of the undoped material (see partial density of states, PDOS, in Fig. 4) and a proton conducting interface (see
Fig. 1), is a sufficiently flexible yet cost efficient supercell. The main features in Fig. 4a are: O$^{2-}$ dominating contribution to the occupied p-states states (red) in the valence band below the Fermi level ($E_F$, dashed vertical line), and Zr$^{IV}$ contributing mainly to the unoccupied states in the conduction band above the Fermi level (green). The band gap of pure ZrO$_2$ is affected only marginally by the hydroxylation; compare Fig. 4a and b. Moreover, in Fig. 4b, new states are introduced at 6–8 eV below $E_F$ owing to the hydroxides. A typical impact on the electronic structure owing to the transition metal ions is shown in Fig. 4c. Here, the presence of Fe$^{III}$ introduces occupied and generally also unoccupied states in the original band gap of ZrO$_2$.

The validity of the model must eventually be assessed on its performance. In a previous study, it referred to in the introduction, connection was made between reaction energies and corrosion properties, see Fig. 3. The apparent validity of the first-principles model was identified by the horizontal dashed black line representing the hydrogen pick-up fraction during oxidation of pure zirconium by water. Moreover, the experimentally observed trends in HPUFs for the suite, Ti, V, Cr, Mn, Fe, Co, Ni, Cu, and Zn, correlate remarkably well with the corresponding energies. A test of the stability of the model is offered by comparing the sensitivity of the energies to the charge and ionic radii of “dummy” ions (Na$^+$, Ca$^{2+}$ or Sc$^{3+}$) embedded in the grain boundary at the site of the transition metal ions to be evaluated. In spite of this harsh test, the seeming correlation between computed hydrogen evolution energies and experimental HPUF for Zr remains.

III. Results and discussion

The correlation between computed hydride–proton recombination energies and observed effective hydrogen pick-up fraction, HPUF, during zirconium oxidation by water implies the existence of a detailed underlying reaction mechanism; this is revealed below. First, structural support for the hydride–proton recombination reaction channel is provided. Second, the origin of the element dependent reaction energy is addressed. Third, properties of the hydride reaction channel are understood to favour the effective electrostatic potential at the reaction site. Support for this conjecture is provided in Fig. 5 and Table 1, which show how atomic structures depend on the choice of “dummy” ion, Na$^+$, Ca$^{2+}$ or Sc$^{3+}$.

For the three “dummy” elements, an intermolecular hydride–proton bond is inferred i.e. a Zr$^{IV}$–H$^+$–OH–Zr$^{IV}$ moiety. In Fig. 5 this is shown in the overlapping partial densities of states of hydrogen belonging to hydride and hydroxide moieties. Seemingly, both the reaction energies and the distances between the hydride and the proton are affected by the charges of the proximal “dummy” ions, see Table 1. The distance between the hydride and proton decreases with reduced charge, suggesting effective lower attraction between the hydride and “dummy” ion with reduced ionic charge in conjunction with less repulsion between the “dummy” ion and the proton. If the “dummy” ion is removed altogether, no local minimum on the potential energy surface corresponding to the hydride–proton structure exists.
H₂ evolution. This relaxation energy depends in a complex way on ionic radius, charge, and change in chemical bonding to the surrounding oxygen ions (mainly for Sc³⁺).

The enthalpy drive for any chemical reaction, including the hydrogen evolution reaction, is found in the favourable bonding of the products as compared to the reactants. Hence, the determination of a structural change, driving the hydride–proton recombination, is an important result of the present study. Indeed, the origin of the shape of the graph displayed in Fig. 3 is associated with the changes in the coordination of transition metal ions upon release of H₂ into the grain boundary. In order to see this it becomes helpful to note that most of the transition metal ions investigated here are 6-coordinated. The chemical changes upon HER are associated with the 6th coordination to TM²⁺/TM³⁺; it is a hydride ion in the reactant and an oxide ion in the product, see Fig. 6a–d. When elementary concepts of ligand field theory are used for the quasi-octahedral systems, the differences become controlled by the occupation of e_g. Consider that, rather than being released, the H₂ molecule may be viewed as squeezed into the grain boundary by the resulting 6th TM–O²⁻ coordination. When the incoming oxide lone-pair encounters an occupied matching TM 3d orbital, the energy gains may be small or even negative. However, if this 3d orbital is empty, then the reaction becomes highly favoured. This interpretation is consistent with the overall trend in Fig. 3. Hence, significant stabilizations are obtained when at least one e_g component is empty, Ti³⁺, V³⁺, Cr²⁺, and low-spin Fe³⁺ for example. The crucial impact that an unoccupied e_g component has on the resulting energy change is clarified in the comparison between low-spin Fe³⁺ and high-spin Ni³⁺, Fig. 3. For these closely neighbouring ions in the periodic table, this effect is reflected in the structural differences of resulting TM²⁺/O²⁻ coordinations, see Fig. 6b and d.

This interpretation is further supported by comparing the similar energy changes of high-spin Mn²⁺ with high-spin Fe³⁺ (red dot in Fig. 3), both of which have two unpaired electrons in e_g. The semi-quantitative validity of the argument may be more fully appreciated by comparing these two systems to the result for low-spin Fe³⁺.

### Table 1

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<th>Distance [Å]</th>
<th>Energy [eV]</th>
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<tr>
<td>Na⁺</td>
<td>1.173</td>
</tr>
<tr>
<td>Ca²⁺</td>
<td>1.262</td>
</tr>
<tr>
<td>Sc³⁺</td>
<td>1.372</td>
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Instead H₂ is formed. Interestingly, no simple correlation between ion charge and reaction energy is seen, see Table 1. This is because the reaction energy is determined to a great extent by the structural relaxations of the products, i.e. subsequent to the H₂ evolution. This relaxation energy depends in a complex way on ionic radius, charge, and change in chemical bonding to the surrounding oxygen ions (mainly for Sc³⁺).

Finally, the impacts of the “dummy” ions on the H₂ evolution reaction energies are shown in Fig. 3 where the black dashed line represents the HPUF for pure zirconium. The three thin blue dashed lines represent hydride–proton recombination energies for the three “dummy” atoms. It is gratifying to note the limited variations in energies, marked by a green shaded band, in spite of the large differences in ionic charges. This offers a stability test for the proposed correlation between the HPUF and reaction energy of the computational methodology.

### III.b. Structural origin of hydride–proton recombination energy

The enthalpy drive for any chemical reaction, including the hydrogen evolution reaction, is found in the favourable bonding

![Fig. 5](image-url)
In particular, the net endothermic hydrogen evolution reactions for the hydrides of Ni$^{II}$ and Co$^{II}$ are due to the insufficient stabilization of TM by the new coordination to O$^{2-}$ as H$_2$ is released. The smallness of this stabilization is due to the Pauli repulsion between the occupied 3d$^5$ orbital and the incoming O$^{2-}$ lone-pair replacing the hydride. Interestingly, with Co$^{II}$ it is the low-spin state, which emerges as the ground state. The low-spin 3d$^7$ electron configuration implies that one e$_g^*$ is vacant. However, it can be seen that both hydride and the incoming O$^{2-}$ lone-pair interact mainly with the occupied e$_g^*$ component due to a Jahn–Teller distortion. Two other exceptions to the emerging understanding are Cu$^{II}$ and Zn$^{II}$. However, the impact of the occupation of e$_g^*$ is expected to be reduced beyond high-spin Ni$^{II}$ if one takes into account the effect of the shrinking of the ionic radius upon an increasing atomic number for elements with the same formal charge and belonging to the same period in the periodic table.

Viewing this result from the perspective of catalysis, a slight endothermicity for hydrogen evolution would indeed be an ideal property for a catalyst. However, Ni$^{II}$ and Co$^{II}$ are reluctant to release H$_2$ into the confining metal/oxide interface; this makes Ni$^{II}$, which has been used extensively in zirconium based nuclear fuel claddings, a “Trojan horse” for hydrogen pick-up.

**III.c. Mechanism for hydride–proton recombination**

The hydride–proton recombination reaction, evaluated in the present study, postulates the release of H$_2$ into a confining hydroxylated ZrO$_2$ grain boundary. The impact of this confinement is to resist the HER. To pass from the hydride–proton reactants to the corresponding products (as described in Section III.b., see [R3]) two reaction pathways emerge, differing by the presence or absence of a TM–H$_2$ intermediate. Here, this property offers a complementary perspective, to that given in Section III.b., on the interpretation of the volcano shape of Fig. 3. The stabilities of intermediates relative to their respective reactants and products are shown for a few of the systems investigated (Fig. 7).

In unravelling the origin of the disparate behaviours of the different systems, we employ as one important component the same elementary concepts of ligand field theory as used in Section III.b. Hence, we note that low-spin Fe$^{III}$ and Zn$^{II}$ display similar reaction energy changes, but what differs is the presence of an Fe$^{II}$–H$_2$ intermediate. The structural characteristics of this intermediate can be appreciated by looking at Fig. 8d and Table 2. Thus, the Fe$^{II}$–H$_2$ moiety has replaced the 6$^{th}$ Fe$^{III}$ coordination to the hydride by the one to H$_2$, compare Fig. 6a and d. The distinct elongation of the H–H bond, see Table 2, is taken to infer that the 1s$^2$ electron pair of H$_2$ becomes shared with the empty e$_g^*$ component of Fe$^{III}$ in a three-centre dative bond. The universality of this TM–H$_2$ intermediate was explored by considering the strongly Jahn–Teller distorted low-spin Ni$^{II}$ (see Fig. 8e and f), which has one doubly occupied e$_g^*$ orbital and one that coordinates the H$_2$ molecule. Indeed, similar bonding was observed for low-spin Fe$^{II}$–H$_2$ (see Table 2). The increased elongation of the H–H bond suggests an even stronger dative bonding for low-spin Fe$^{III}$ than for low-spin Fe$^{II}$. This is believed to be due to a larger field acting on H$_2$ for low-spin Ni$^{II}$ due to the smaller ionic radius. The inability of 3d$^{10}$ Zn$^{II}$ to provide sufficient stabilization of TM by the new coordination to O$^{2-}$ for Mn$^{II}$ with the empty e$_g^*$ orbital and the lone-pair of H$_2$ becomes shared with this ion. To emphasize the difference between occupied and unoccupied e$_g^*$ orbitals further, we note that neither high-spin 3d$^6$ Fe$^{II}$ nor high-spin 3d$^5$ Mn$^{II}$ possess any TM–H$_2$ local minimum along the reaction coordinate considered.

An apparent exception to the rule is provided by high-spin 3d$^8$ Ni$^{II}$–H$_2$ (see Fig. 8b and d). However, when the H–H bond distance was inspected, only a slight deviation from that of free H$_2$ was found (see Table 1). Rather than dative bonding between TM and H$_2$, the molecule remains at the TM site because the oxy-hydroxide ligands do not provide sufficient power to cause the H$_2$ molecule to detach into the confining hydroxylated.
This is consistent with high-spin NiII displaying the longest TM–H distances, which corresponds to the weakest Coulomb field. These findings are consistent with the resulting endothermicity for the hydrogen evolution reaction (HER) for high-spin NiII (see Fig. 7); while both high- and low-spin NiII display endothermicity for the HER, the natures of the NiII–H2 moieties are qualitatively different.

Finally, the analogy between low-spin FeII and low-spin NiII is further substantiated by comparing the corresponding energy changes during TM–H2 decomposition, whereby H2 is released and the oxygen coordination sphere of the corresponding TMII is completed. In both cases this reaction step becomes exothermic by 0.6 eV (see Fig. 7).

### III.d. Correlation between hydride–proton recombination energetics and HPUF: implications concerning of nickel

The hydride–proton recombination pathway for hydrogen evolution having been mapped, a connection is made between the experimental observations and the emerging mechanistic interpretation. The correlation observed between HPUF and reaction energy changes, rather than activation energies, suggests the HPUE to be essentially of thermodynamic origin rather than due to kinetics. Thus, instead of competing reaction barriers, the decisive effects of various transition metal ions on the overall hydrogen evolution reaction are determined by the stabilities of the grain boundary hydride intermediates relative to the H2 molecule released into the grain boundary. Let the effective potential drop be \( \sim 1.0 \text{ eV per } \text{H}_2 \). This number is taken from the fact that TiIII decorated grain boundaries display negligible HPUE (see Fig. 3). Thus, we arrive at the residual net drive, \( U \), for hydrogen evolution inside the hydroxylated interface:

\[
U \approx 1.0 \text{ eV}/2e^- = 0.50 \text{ V}. \tag{1}
\]

Now, let the charge associated with the oxide formation due to hydrogen evolution and with the hydrogen pick-up, be \( Q_{HE} \) and \( Q_{HPU} \), respectively:

\[
Q_{tot} = Q_{HPU} + Q_{HE}. \tag{2}
\]

From the definition, we have

\[
\text{HPUE} = \frac{Q_{HPU}}{Q_{tot}} = 1 - \frac{Q_{HE}}{Q_{tot}} = 1 - \text{HEF} \tag{3}
\]

where HEF, hydrogen evolution fraction, is the fraction of water associated protons which are converted to H2. Eqn (3) allows us to arrive at the observed correlation by acknowledging that

\[
\text{HEF} = \frac{Q_{HE}}{Q_{tot}} = \frac{UQ_{HE}}{UQ_{tot}} = \frac{\Delta E_{HE}}{\Delta E_{tot}}. \tag{4}
\]

Having determined the correlation between HPUE and the energy changes for hydride–proton recombination, the detrimental impacts of NiII or CoII in the grain boundary may now be understood more fully. Indeed, the harm caused by these ions is dual, in that besides reluctance to release H2 (Fig. 3 and 7), a “back-door” mechanism for HPU is also provided. Hence, complementarily to the endothermicity of the hydride–proton recombination energy changes for NiII or CoII associated hydrides, a priori CoII or NiII decorated grain boundaries display an affinity to any H2 present in the grain boundary (possibly

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<td>1.682</td>
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interface. This is consistent with high-spin NiIII displaying the longest TM–H distances, which corresponds to the weakest Coulomb field. These findings are consistent with the resulting endothermicity for the hydrogen evolution reaction (HER) for high-spin NiIII (see Fig. 7); while both high- and low-spin NiIII display endothermicity for the HER, the natures of the NiIII–H2 moieties are qualitatively different.

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Fig. 8 Depicted are the hydride reactant structures: (a) for FeII low-spin, (b) for NiII high-spin and (c) for NiII low-spin. Also displayed are the intermediate TM–H2 structures: (d) for FeII low-spin, (e) for NiII high-spin and (f) for NiII low-spin. Energy for the structures can be seen in Fig. 7. FeII and NiII high-spin have an octahedral coordination, while NiII low-spin displays a square planar coordination.

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Table 2 Bond distance within the TM–H2 intermediate structure. The distance between the two hydrogen atoms in the hydrogen molecule in the gas phase is 0.748 Å.

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released from an FeH site) to form an NiII or CoII associated hydride in conjunction with a hydride proton.

This is illustrated in Fig. 9. In step (A)–(C) the detailed dependence of formation of H2 on the transition metal is repeated (Fig. 3 and 7). The subsequent step, (C)–(A), demonstrates how NiII and CoII become sinks for the H2 released.

IV. Concluding remarks

A hydrogen evolution reaction, HER, mechanism in a hydroxylated zirconia grain boundary was identified, by which H2 is formed by hydride–proton recombination. This reaction occurs between hydride associated protons and transition metal associated hydrides. Characteristic intermediates along this reaction coordinate were described. A similar mechanism for hydrogen evolution was recently proposed for a nickel electrocatalyst supported by seven-membered cyclic diphosphine ligands containing one pendant amine, with the Ni supporting the hydride and the amine providing the proton in the hydride–proton recombination reaction. Moreover, while a di-hydride intermediate was postulated for the Ru catalyst dehydrogenation of methanol, it is conceivable that the hydride–proton recombination reaction is also applicable here. Thus, common ground is implied for the modelling of properties relevant to water induced corrosion of zirconium based alloys and of electrochemical (reverse) fuel cells.

In this way, density functional theory was employed to model the impacts of all 3d elements on the cathode process for the water induced oxidation of zirconium alloys. The robustness of the grain boundary model was also assessed. A major finding of the present study is that the energy change of the HER inside a grain boundary was also assessed. A major finding of the present study is that the energy change of the HER is determined by the competition for a site in the coordination

The energy changes of the HER inside a grain boundary were analysed and the release of H2 was found to result from a “squeezing” process. Repeatedly, the electron population in the corresponding e_g was found to be decisive, i.e. both for the energy changes and for the presence of a bound TM–H2 intermediate. Consequently, sensitive dependence of the HER on the choice of spin state is observed when the different spin states cause different e_g occupations.

The correlation observed between hydrogen pick-up fraction and hydrogen evolution energy changes was discussed; it indicates a purely thermodynamic origin. Dual causes were identified for the detrimental impacts of nickel and cobalt on hydrogen pick-up in zirconium alloys.

Acknowledgements

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