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Field effects explain the unintuitive potential response of electrochemical oxygen evolution in acid†

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Electrochemical water oxidation (OER) is the most important electrode reaction in electrocatalysis, representing the default counter-reaction in the plethora of modern electroreductions. Given this prominent role in the electrochemistry-based green transition, improving its efficiency is of utmost importance. Here, identifying novel catalysts by means of computational screening necessitates clarity on the reaction mechanism as this is used to decide on appropriate activity descriptors. To date, however, the mechanism of OER, even on the most widely used catalyst in acid, IrO₂, is still debated, and the debate is fuelled by the consistent appearance of Tafel slopes indicative of a non-electrochemical rate-limiting step. Here, we employ density functional calculations and microkinetic modelling to analyse the mechanism of acidic OER on IrO₂(110), with an emphasis on the polarization of reaction intermediates. Introducing this degree of freedom shows that the electrostatic destabilization of surface-bound oxygen atoms with increasingly positive potentials increases the effective potential response of the reaction. Thus, a reaction mechanism through OOH*⁻-formation could be confused with a non-electrochemical rate-limiting step. Furthermore, we highlight that a mechanism limited by the desorption of adsorbed O₂ is unlikely, as this step is facile at room temperature, but caution is needed in treating adsorbed O₂ in GGA-DFT. By incorporating these elements into our model, we simulate Tafel curves that reproduce the experimental potential response in both the low and high overpotential regions, pointing out that including surface dipole effects is essential for understanding and reproducing experiments.

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Sustainability spotlight

The oxygen evolution reaction (OER) plays a critical role in the development of efficient and sustainable energy solutions, particularly in water splitting for hydrogen production. However, the dependence on trial-and-error experiments has made it difficult to find promising electrocatalysts. By applying DFT calculations and microkinetic modeling to OER, we can pinpoint key intermediates, rate-determining steps, and reaction pathways that are central to improving the catalyst performance. This approach can significantly minimize the discrepancy between theory and experiment, making it easier to identify promising materials and conditions that lead to higher efficiency and lower environmental impact. This aligns with the goals of the creation of energy-efficient technologies for renewable energy production where understanding of the OER mechanism offers a promising route for advancing sustainable energy technologies.

Introduction

The oxygen evolution reaction (OER) is crucial to achieve a fossil fuel-free economy.¹ When powered by renewable energy, OER supports the sustainable production of hydrogen, hydrocarbons, and other valuable chemicals. However, oxidizing potentials and the corrosive environment at the anode limit the efficiency of OER, making it a key area for improvement.

Further, state of the art catalysts (IrO₂ and RuO₂) require significant overpotentials (~300 mV) to function effectively, which adds to energy losses in nearly all electroreduction reactions of interest.¹ The overpotential and stability could be reduced slightly through modifications such as, varying catalyst supports² or introducing alternative metals into IrO₂/RuO₂.³ However, although representing the likely most applied half-reaction in electrocatalytic technology, designing better OER catalysts remains a challenge, even down to the level of knowing the relevant reaction steps, as the mechanism of *e.g.* acidic OER on its historically most used catalyst, IrO₂, is still under debate.

OER requires the transfer of four electrons across the electrode/electrolyte interface and the generation of four protons *via* the stepwise decomposition of water followed by the

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Scheme 1 Schematic illustration of the possible reaction mechanism for OER on IrO₂(110). The O₂ molecule is formed by O–O bond formation either by a Langmuir–Hinshelwood (LH) mechanism or Eley–Rideal (ER) mechanism *via* a coupled proton–electron transfer. Possible rate-limiting steps (RLS) are marked in blue and green colors.

formation of an O–O chemical bond, as shown in Scheme 1. In the 1960s, Bockris *et al.* suggested 14 possible routes for OER on Pt, proposing primary water discharge as the rate-determining step.⁴ Conversely, on RuO₂(110), OER was suggested to occur *via* the Eley–Rideal (ER) mechanism, with the coupling of surface-bound oxygen with H₂O from the solution being reported as the rate-limiting step (RLS).⁵ In contrast, the Langmuir–Hinshelwood (LH) mechanism (such as by coupling two oxygen atoms on the surface) was deemed energetically less favoured and weakly influenced by the applied potential. Finally, a participation of lattice oxygen in the reaction could also be hypothesized (Mars–Van Krevelen mechanism). Kasian *et al.* and Scott *et al.* addressed this possibility by O-isotope labelling of the reacting water and the oxide catalyst, respectively. Both studies concluded that on rutile IrO₂, the vast majority of oxygen is formed out of interfacial water.^{6,7} While lattice oxygen could be a minor contributor in RuO₂ and IrO₂, it can be important in other materials, such as perovskite (*e.g.* SrIrO₃,⁸ LaCoO₃,⁹) and Co-based catalysts (*e.g.* Co₃O₄).¹⁰

Computational analysis of the thermodynamic landscape in OER reveals that the formation of OOH* is the most difficult step for many electrocatalysts including IrO₂,^{11,12} and a further analysis by Dickens *et al.*, including explicit calculations of potential-dependent activation energies confirmed that O–OH bond formation is most likely the RLS in all cases.¹³ However, the O–O coupling pathway *via* a LH mechanism, involving a chemical RLS between two oxygen atoms, was suggested in earlier studies due to OER's potential response above ~1.5 V *vs.* RHE.^{14–16} Additionally, S. Kwon *et al.* concluded that the OER mechanism and RLS depend on the facet and potential of IrO₂.¹⁷ They concluded that direct oxygen coupling is favourable on the (101)-facet of IrO₂, while the other facets follow a mechanism limited by the nucleophilic attack of water.

Recent theoretical analyses further suggested that, at high overpotentials on IrO₂, the final oxygen desorption step may become thermodynamically unfavourable.^{18,19} As an alternative to direct O₂-desorption from the catalyst surface, recent

findings by Binninger *et al.* suggested that OER on IrO₂ may follow the association of two O–O* species, especially at high overpotentials.²⁰ Exner further validated this result^{21,22} concluding that the *OO–OO* recombination mechanism is crucial and should not be overlooked in future screening studies of OER.

All these deviating mechanistic interpretations need clarification to justify specific catalyst screening strategies. Turning to experiments, one of the strongest mechanistic tools in electrocatalysis for a distinction between the varying pathways is the interpretation of the potential needed to change the current by one order of magnitude, the so-called Tafel slope. From empirical models, such as Butler–Volmer theory in the absence of mass transport limitations,²³ the Tafel slope is a consequence of the transfer coefficient α following the relationship:

$$\alpha = \frac{k_B T}{e} \ln(10) \frac{\partial \log_{10}(j/j_0)}{\partial U} \approx \frac{59 \text{ mV}}{\text{Tafel slope}}, \quad (1)$$

with $\frac{\partial \log_{10}(j/j_0)}{\partial U}$ referring to the change of current density j with applied potential U and k_B , T , and e representing the Boltzmann constant, temperature, and the charge per electron, respectively. Fig. 1 presents a collection of experimental data for OER on IrO_x-based catalysts in acid, including both films and nanoparticles (NPs).^{24–31} Although few data span a broad potential window, for those cases, it is evident that OER exhibits two distinct Tafel slope regions. The potential at which this transition occurs defines the low and high-potential Tafel regions for the data below and above, respectively. The low-potential Tafel region slope is about 40 mV dec⁻¹, representing $\alpha \approx 1.5$. However, the more commonly investigated (and commercially more relevant) high-potential region exhibits Tafel slopes of 60–72 mV dec⁻¹ (α between 0.83 and 1) for films and around 50 mV dec⁻¹ for NPs ($\alpha \approx 1.2$). Previous theoretical studies successfully reproduced low-potential region behaviours and overestimated values for higher potential ones,¹³



coupled electron transfer (PCET)-based RLS at low overpotential ($\beta_{\text{RLS}} < 0$), to a RLS defined by a potential-independent chemical step ($\beta_{\text{RLS}} = 0$).

The picture of expected Tafel slopes changes once the system is allowed to deviate from the idealized values. Two deviations can be envisioned; pinning the value of β_{RLS} to -0.5 for PCET steps or zero for formally non-electrochemical steps has no strong justification, where β_{RLS} can, in principle, adopt any value between the effective potential responses of the initial state and final state of the elementary steps.^{37,38} Further, neglecting the contribution of the polarization of reaction intermediates, γ ,³⁴ is a crude approximation. Here, the field response of the RS, γ_{RS} , is particularly important, as it will directly influence $\frac{\partial \Delta G_{\text{act}}}{\partial eU}$. While often negligible, particularly the binding of oxygen-containing adsorbates can exhibit a substantial inherent dipole, which changes α into:

$$\alpha = N_{\text{pe}}^{\ddagger} - N_{\text{pe,RS}} - \beta_{\text{RLS}} + \gamma_{\text{RS}} \quad (6)$$

Note that, analogous to β_{RLS} , we chose the sign of γ_{RS} to be consistent with $\left(\frac{\partial G_{\text{RS}}}{\partial eU}\right)_{\text{PH}}$ rather than α .

Fig. 2 shows the variations in the Tafel slope, resulting from variations in γ_{RS} and β_{RLS} . We highlight the combination of Tafel slopes for RS conditions of a monolayer of OH* (y-axis) and O* (x-axis) defining the RS of the catalyst. The two cases differ in $N_{\text{pe}}^{\ddagger} - N_{\text{pe,RS}}$, being 1 and 0 for OH* and O* coverage, respectively. This difference results from the need of OH* to traverse an O*-intermediate to reach the RLS, while O* does not have an equilibrated elementary step preceding the RLS.

Fig. 2a focuses on the mechanism involving O–OH* formation as the RLS. Here, $\beta_{\text{RLS}} = -0.5$ together with γ_{RLS} of both OH* and O* being zero (light turquoise line, $\gamma_{\text{OH}} - \gamma_{\text{O}} = 0$) results in the expected combination of 120 and 40 mV dec⁻¹ on the x and y-axes, respectively. While keeping β_{RLS} constant,

increasing γ_{OH} and γ_{O} results in a reduction of their respective Tafel slopes, where e.g. $\beta_{\text{RLS}} = -0.5$, $\gamma_{\text{O}} = 0.2$ and $\gamma_{\text{OH}} = 0.1$ lead to Tafel slopes of 86 and 38 mV dec⁻¹.

The dependency of the Tafel slopes on γ_{RS} in the LH mechanism, shown in Fig. 2b, is even more prominent as its effect is doubled due to the participation of two O*-species in the RLS. Equally, $N_{\text{pe}}^{\ddagger} - N_{\text{pe,RS}}$ in this case takes the values of 2 and 0, for OH* and O* covered surfaces, respectively. Note that, in this mechanism, $\beta_{\text{RLS}} \rightarrow 0$ is likely, making the potential response purely defined by γ_{RS} and leading to large Tafel slopes in the high overpotential region.

Overall, Fig. 2 shows that γ_{RS} is a central component of the measured potential response and leads to a large variety of possible Tafel slope combinations in OER, independent of the actual mechanism at play.

To estimate the magnitude of γ_{RS} , in Fig. 3 we show the DFT-predicted response of the differential formation free energy of the relevant intermediates of OER to an applied homogeneous electric field perpendicular to the electrode surface at a full background coverage of OH* (Fig. 3a) and O* (Fig. 3b) on the CUS sites of IrO₂(110) (see ESI Section 1† for Computational details). Analogous results for the field dependence of the change of surface phases by means of average formation free energies are shown in ESI Section 3,† with nearly identical values.

We found that, in the vicinity of the potential of zero charge (PZC) of the electrode, O*, OH*, and O₂* are destabilized by negative field strengths, corresponding to more anodic conditions, while H₂O* and OOH* are stabilized. This varying behaviour results from the varying surface dipole changes upon adsorption, $\Delta\mu$, and, thus, γ , as summarized in Fig. 3c. We provide the framework to define γ from the change in surface dipole $\Delta\mu$ and polarizability $\Delta\eta$ in ESI Section 4.†

γ_{O} is generally enhanced relative to γ_{OH} , which can be attributed to its binding motif in a top site on the IrO₂(110) CUS, forming a distinct dipole perpendicular to the catalyst surface.³⁹



Fig. 2 Tafel slopes considering (a) the ER mechanism (O–OH formation) and (b) the LH mechanism (O–O coupling). Contours show the variation in the Tafel slope considering the potential response (γ) of OH* and O* for estimating low (y-axis) and high (x-axis) Tafel slopes, respectively. Experimentally observed Tafel slopes are marked with a dark green box.





Fig. 3 Field response of the relevant intermediates in OER on IrO₂(110) under (a) OH* coverage and (b) O* coverage on Ir_{CUS} sites. (c) Table with fitted values for the change in surface dipole ($\Delta\mu$, [eÅ]) and polarizability ($\Delta\eta$, [$\frac{\text{e}\text{\AA}^2}{\text{V}}$]) associated with a change in surface phase. Additionally, the net potential response for each adsorbate is indicated where $\tilde{U} = U_{\text{SHE}} - U_{\text{PZC}}$.

Conversely, OH* tends to form a hydrogen bond network with the hydrogen pointing parallel to the surface, reducing its overall dipole perpendicular to the surface.

We now show how γ , and particularly its relative differences influence phase stabilities. Fig. 4a shows the calculated Pourbaix diagrams for IrO₂(110) including interfacial field effects (see ESI Section 5† for calculation details). In agreement with previous results,³² we identified that IrO₂(110) is covered by OH* on the IrO₂(110) CUS when ramping up the potential to 1.52 V vs. RHE, where it transitions into a monolayer of O*. We also identified that the deprotonation of the bridge-O (O_b) is likely responsible for the capacitive peaks in cyclovoltammetry (CV) appearing at intermediate positive RHE potential, as reported in ref. 26 for single-crystal IrO₂(110). This phenomenon has also been documented in amorphous IrO_x and rutile IrO₂ in acid.^{31,32}

The field response of the varying surface phases has a minor effect on the transition from OH*- to O*-covered IrO₂(110) CUS sites when compared to the case without the inclusion of the field effect shown in ESI Sections 6 & 7.† Conversely, OH* formation on IrO₂(110) CUS shifts by ~ -0.7 V. This behavior is a consequence of our choice of PZC in the field response model,

which we chose based on the calculated value for the bare IrO₂ surface (see Fig. S2†). As the OH*- to O*-covered IrO₂(110) CUS-transition happens close to the PZC, the field is small and thus has little influence on it. We note that the PZC does influence the specific transition potential, but its change with potential, our central benchmark, is less affected.

Differences in γ of the two phases involved in the transition are expressed as pH-dependence on the RHE potential scale,³⁴ which allows us to directly benchmark our determined values against experimental observations. In Fig. 4b, we benchmarked the potential where the RS changes from a monolayer of OH* to a monolayer of O* on the CUS of IrO₂ against experimental validations from Kuo *et al.*, who studied the transition between the two phases based on the appearance of oxidation/reduction waves in CVs.²⁶ The fingerprint CVs of IrO₂(110) were recorded in a wide pH range, exhibiting a non-Nernstian shift in the peak position of two phase transitions between 0.8 and 1.6 V vs. RHE. Based on Fig. 4a, we can attribute the two waves to the electrochemical desorption of H* from the O_b sites (0.87 V vs. RHE) and the transition from OH* to O* (1.52 V vs. RHE) on IrO₂(110) CUS. With increasing pH, the experimental peak positions





Fig. 4 Surface phase diagram for different coverage conditions on the pristine $\text{IrO}_2(110)$ surface at the full coverage limit with field effects (a). Potential vs. RHE where full OH^* coverage transitions to full O^* coverage at different pH when no field effect is included (white circles), when a field effect is included (solid black circles) and from experimental data (black triangles)²⁶ (b). $^*\text{Ir}_{\text{CUS}}$ represents empty Ir_{CUS} sites and O_b an oxygen bridge.

shifted in the negative direction against the RHE potential scale, which allows us to directly probe the γ_{O} with respect to γ_{OH} . The experimental behaviour is in line with the results from our calculations including the calculated γ_i , with $\frac{\partial U_{\text{RHE},x}}{\partial \text{pH}} = -10.9 \text{ mV pH}^{-1}$. This proves the qualitative behavior that O^* is destabilized by increasingly anodic potentials more than OH^* , strongly indicating that γ_{O} may substantially influence on the Tafel slopes in the high overpotential region. Spin polarized results for the same are shown in Fig. S3.†

After having identified the relevant surface phases at reaction conditions and estimating γ_{OH} and γ_{O} , we now show how both these aspects influence experimental OER observations. For this, Fig. 5 shows the calculated free energy diagram at pH zero for all the intermediates involved in the (a) ER and (b) LH pathways. The effective changes of the free energy of each state with U at constant pH, $\left(\frac{\partial \Delta G_i}{\partial eU}\right)_{\text{pH}} = N_{\text{pe},i} + \gamma_i$, are explicitly given. For clarity, only the energetics on an O^* -covered surface are shown, while corresponding energetics with a full OH^* coverage as the RS are shown in Fig. S6.† and the qualitative picture is equal in both cases. The direct link to experimental observations is made in the corresponding lower panels (c and d) of Fig. 5 resulting from a microkinetic model (see section 8 in ESI† for more detail).

Starting from an empty $\text{IrO}_2(110)$ CUS-site, the adsorption of water to form H_2O^* does not involve a proton–electron pair

($N_{\text{pe},^*\text{H}_2\text{O}} = 0$). However, H_2O^* is considerably stabilized by increasing the potential, and when calculating $\gamma_{\text{H}_2\text{O}}$ at 1.23, 1.43, and 1.63 vs. RHE the average value is $\gamma_{\text{H}_2\text{O}} = -0.29$, defining its whole potential response. Next, one proton–electron pair is produced for reaching OH^* ($N_{\text{pe},^*\text{OH}} = 1$). Combined with γ_{OH} , the effective potential response of OH^* formation resulted in -0.98 eV/V . Subsequently, to produce O^* , two proton–electron pairs are created with respect to the empty site ($N_{\text{pe},^*\text{O}} = 2$). However, the considerable γ_{O} partially counters the purely Nernstian potential response leading to an effective $\frac{\partial \Delta G_{^*\text{O}}}{\partial U} = -1.80 \frac{\text{eV}}{\text{V}}$. O^* is followed by the RLS in both considered reaction mechanisms. For the ER mechanism shown in panel (a), we used the activation energy and $\beta_{\text{O-OH}} = -0.58$ as determined in our previous work.¹³ Here, Dickens *et al.* calculated the activation energy for the formation of OOH^* on oxides and investigated its dependence on the applied potential (See ESI Section 8† for further details). $N_{\text{pe},^*\text{O-H}_2\text{O}} = N_{\text{pe},^*\text{O}} = 2$, as no extra proton or OH^- are involved for reaching the transition state under acidic conditions. After crossing the RLS barrier in the ER mechanism, OOH^* is reached, which, in contrast to O^* and OH^* , is stabilized by an increase in the negative field (more positive potential), resulting in an enhanced potential response compared to the field free case (*i.e.* $\alpha_{\text{OOH}^*} - \alpha_{\text{O}^*} = 1.71$).

Finally, O_2^* results from the deprotonation of OOH^* , which exhibits a net potential response of -3.93 eV/V . Note that we did not use the bare DFT calculated formation free energy of O_2^* in





Fig. 5 Free energy diagrams and simulated current densities for OER including effects of the electric field on IrO₂(110). Panels (a) and (b) show free energy diagrams for the ER and LH mechanisms, respectively, under full O* coverage. The mean values of the effective changes of the free energy of each state with U at constant pH, $\left(\frac{\partial \Delta G_i}{\partial eU}\right)_{\text{pH}}$, are noted for each step. Panels (c) and (d) show the simulated current density (j) at pH = 0 for the ER and the LH pathways, respectively. The solid lines represent the effective current taking into account the change in surface phase. Coverages obtained from the microkinetic model are shown in blue for OH (θ_{OH}) and O (θ_{O}). The obtained Tafel slopes for the pre-kink and post-kink regions are given in panels (c) and (d).

the free energy diagram in Fig. 5, as it is overstabilized by the applied GGA-DFT methodology. Instead, we used a value benchmarked by employing the Redhead analysis of experimental temperature-programmed profiles (TPD) of O₂ desorption from IrO₂(110).^{40–42} We describe the process of determining the energy in more detail in ESI Section 9† and will discuss the need for this correction for a correct mechanistic analysis in the discussion of the microkinetic results below.

For the definition of α in the ER mechanism (α_{ER}), we can invoke eqn (6), hence simply calculating the potential response between the RLS transition state (-2.58 eV/V) and the RS, *i.e.* -0.98 eV/V and -1.80 eV/V at low and high overpotentials, respectively. Thus, the predicted α_{ER} values are 1.6 and 0.78. In the LH mechanism, shown in Fig. 5b, 2O* species act as the reactants of the RLS. Thus, the energy landscape is altered involving two adsorbates throughout up to the RLS. Analogously, the potential response of each step before the RLS is doubled as well. As the RLS represents a formally non-electrochemical step in this mechanism, the potential response between 2O* and O₂(g) only varies by 0.4 eV/V, purely defined by the quenching of γ_{O} upon desorption. Thus, α_{LH} calculated from Fig. 5b results in 2.04 and 0.4 for the low and high overpotential regions, respectively.

A comparison of the simulated polarization curves in Fig. 5c and d reveals that the ER mechanism aligns much more closely

with experimental data both in terms of Tafel slopes and absolute current densities, which approach at 1 mA cm^{-2} at ~ 1.6 V. Conversely, the LH mechanism, exhibits negligible reaction rates throughout the potential range because of the huge thermodynamic barrier connected to the formation of O₂ from two O* species.

As expected, α_{ER} directly translates into the Tafel slope estimates of 37 mV dec^{-1} and 77 mV dec^{-1} for the low and high overpotential regions, respectively, under O* coverage (see Fig. 5c). Note that this result is only obtained when the O₂* to O₂(g) energy value is set to ~ 0.31 eV, as fitted from the TPD analysis shown in Fig. S7,† as O–OH* formation remains the RLS within the simulated potential range. Specifically, including the field effect for O₂* shows that this species destabilizes at oxidative potentials, weakening its interaction with the surface compared with pure TPD-derived data, then when the potential increases, the O₂* to O₂(g) value is even smaller than 0.31 eV. However, this picture changes when using the desorption energy predicted by GGA-DFT (~ 0.81 eV). As the activation energy for O–OH* formation is potential-dependent, it decreases as the potential increases. At a certain potential, it becomes smaller than the desorption energy of O₂*, as predicted by GGA-DFT, making the latter the apparent RLS. This shift alters the Tafel curve behaviour, resulting in a near complete loss of potential response, in strict disagreement with experiments.



In contrast to the ER reaction path, the simulated LH mechanism exhibited Tafel slopes of 29 and 150 mV dec⁻¹ (see Fig. 5d), which also disagrees with the measurements shown in Fig. 1. A degree of rate control analysis (*cf.* Fig. S8 & S9 in ESI†) for this mechanism shows that the chemical O–O coupling step and the energy of O₂* do not influence the overall rate of the reaction, as O₂(g) defines the highest barrier to the product starting from the RS.

Note that we have chosen constant values of C_H (*i.e.* 25 μF cm⁻²) and U_{pzc} (1.5 V vs. SHE) in our model. These choices directly affect the quantitative values of the Tafel slopes and the potentials of surface phase changes. Therefore, we also probed the sensitivity of simulated Tafel plots to variations of C_H (Fig. S10†) and U_{pzc} (Fig. S11 and S12†). The results show that the trends in Tafel slopes are mostly unaffected by the choice of parameters. That is, while the low overpotential region (*OH-covered surface) is virtually unaffected by the interfacial field, the high overpotential region (*O-covered surface) shows a substantial field response, leading to reduced Tafel slopes compared to the case without the field effects.

While in the present study, we focussed only on the most stable (110)-surface of IrO₂, we are confident that analogous characteristic field interactions are present on other facets. Further, the stronger binding of *O on less coordinated surface sites likely even reduces the probability of the LH mechanism. However, specific evaluations on other facets would be needed to confirm these hypotheses.

Overall, our analysis shows that while the proximity of experimental Tafel slopes to the characteristic value of 60 mV dec⁻¹ might make it tempting to assign a chemical RLS. This can be equally well explained by the combined potential response of the RLS and RS in a PCET-limited step, as is the case for the ER mechanism. Conversely, the LH mechanism would never have a value approaching 60 mV dec⁻¹, as it would formally (from purely Nernstian behaviour) only exhibit Tafel slopes of ~30 mV dec⁻¹ or infinity. However, including γ_O leads to non-zero potential responses, which still deviate strongly from the experimentally observed values.

This *a priori* unintuitive Tafel behaviour likely extends beyond IrO₂(110), both in terms of alternative surface facets and other rutile oxides. To account for the behaviour, a further simplified model directly applicable in large scale screening studies can be applied: conventional DFT calculations for the relevant reaction intermediates directly provide μ from the ground state charge density (mostly given in the output of DFT codes) and the work function as an estimate for U_{pzc}. Utilizing them allows a first order estimate of relevant γ values without the need for field dependent (or even more sophisticated) calculations.

Conclusions

In this study, we used DFT calculations and microkinetic modelling to analyse the mechanism of acidic OER on IrO₂(110) including the polarization of reaction intermediates. We have shown that the ER type mechanism through a coupled proton–electron transfer RLS to form OOH* suggested by Dickens

*et al.*¹³ does not contradict the experimentally measured Tafel slopes as long as the field response of reaction intermediates is considered. Further, our results suggest that the combination of two oxygen atoms from the surface *via* the LH mechanism is unlikely, as it would result in negligible current densities that would be marginally affected by the increase in overpotential. Additionally, by analysing experimental data of temperature-programmed desorption profiles, we emphasized that the desorption of O₂* is fast. Hence, it is unlikely to be a RLS. However, GGA-DFT poorly captures this, overestimating the O₂* adsorption energies.

Data availability

All the data for DFT calculated energies are summarized in the ESI.† Data such as inputs for the figures and the atomic structures will be made available at https://github.com/CatTheoryDTU/OER_mechanism_IrO2_acid upon acceptance of the article.

Conflicts of interest

There are no conflicts of interest to declare.

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