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Bypassing the scaling relations in oxygen electrocatalysis with geometry-adaptive catalysts

This communication introduces the concept of geometry-adaptive electrocatalysis, where a catalyst adjusts its geometry during the reaction. A model system of metal-nitrogen-carbon (M-N-C) catalysts – the dual-atom site 2Co-N_4 of variable curvature – proves the concept from the first principles. Density functional theory calculations show how cycling the curvature effect with a geometry adaptation bypasses the scaling relations. Thus, in theory, geometry-adaptive electrocatalysis offers a promising direction to address the current stagnation in the experimentally measured overpotential for oxygen evolution and reduction reactions. It also indicates the possibility of discovering the ideal oxygen electrocatalyst.

Introduction

The "holy grail" of electrocatalysis is the ideal catalyst concept – catalysts that accelerate "dream reactions", such as oxygen evolution and reduction reactions (OER and ORR), using only the energy set by their thermodynamic potential. While the quest for better catalysts motivates much of electrocatalytic research, the ultimate goal of this journey is the discovery of the ideal catalysts. Such catalysts promise 100% efficient OER and ORR for sustainable economic development.

The journey towards the ideal catalysts is theoretically and experimentally challenging due to a fundamental constraint – linear scaling relationships – tying adsorption energy values to the chemical nature, valence, and coordination of reaction intermediates. ^{1,2} In oxygen electrocatalysis, the scaling relation dictates the relative adsorption energies of two key intermediates, OH and OOH. ^{3,4} It is worth noting that in electrocatalysis theory, the adsorption energy (ΔG) serves as the primary descriptor. ⁵ Ideally, the difference between ΔG for OH and OOH intermediates should be 2.46 eV. ⁵ Yet, in

computational practice, the OH–OOH scaling relation stands as: $\Delta G_{\rm OOH} \approx \Delta G_{\rm OH} + 3.20$ eV.⁶ This implies that the OH and OOH intermediates cannot have the ideal ΔG difference.⁷ Furthermore, the deviation from the reaction thermodynamic potential on the $\Delta G_{\rm OH}$ scale follows the Sabatier principle to form activity volcanoes shown in Fig. 1(a).⁷ Specifically, the solid magenta line in Fig. 1(a) highlights the apex of the "associative mechanism" volcano defined by the OH–OOH scaling relation. The predictions from Fig. 1(a) align with experimental potential values in Fig. 1(b), where they converge towards the theoretical limit.

Approaching the theoretical limit in oxygen electrocatalysis calls for creativity in exploring novel catalysts and mechanisms. Historically, platinum group metal catalysts have showcased superior activity, yet they raise valid concerns regarding sustainability and cost. Heal- and nitrogen-doped carbon (M–N–C) materials, known for their single-atom sites (M–N $_x$), have become a more practical alternative thanks to their high catalytic activity. Still, efforts to enhance the catalytic performance of both platinum group metal and M–N–C catalysts consistently encounter the constraint of the OH–OOH scaling relation, as illustrated in Fig. 1(b).

To implement the ideal catalyst, it is essential to find a way to circumvent the OH-OOH scaling relation.³⁷ One strategy is to sidestep the associative mechanism (involving OOH) by introducing an adjacent second active site.³⁸ In this scenario, electrocatalysis can proceed via a dissociative mechanism, substituting the OOH intermediate with O on one site and OH on the other. 10,39 In competition between the associative and dissociative mechanisms, only a few model catalysts, like Fe₂N₆, Co₂N₆, and FeCoN₆, favour the dissociative mechanism. 40,41 The preference depends on the active site metal and coordination. 42 Specifically for the 2Co-N₄ model, the dissociative mechanism is preferred. This results in the OH-O/OH scaling marked with a dot-dashed green line in Fig. 1(a).9 The "dissociative mechanism" volcano suggests a feasible improvement in the ORR as well as enhanced bifunctionality, which is the difference between

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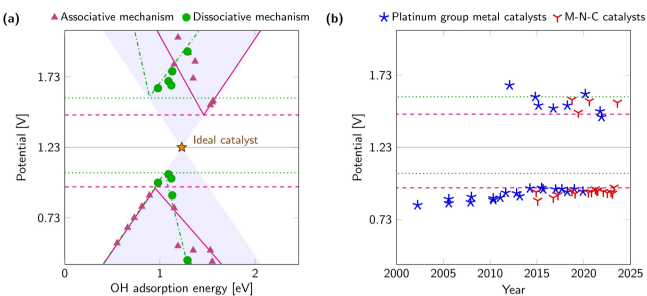


Fig. 1 (a) Overpotential volcano projected on the ΔG_{OH} description. Triangles denote the DFT-calculated overpotentials for single-site M-N-C catalysts, whereas circles represent those for dual-site M-N-C with curvature. Dashed and dotted lines highlight the apex of these overpotential volcanoes. (b) Timeline with outstanding, experimentally measured ORR and OER potentials for both platinum-group metals and metal-carbonnitrogen (M-N-C) catalysts. The selected potentials correspond to a current density of 10 mA cm⁻² for OER and 3 mA cm⁻² for ORR. Data obtained from ref. 10-33.

ORR and OER potentials for a given material. Another intriguing strategy involves altering the catalyst's geometry, as demonstrated by the geometry adaptation of Pt nanoparticles to vary the intermediates' adsorption energy.⁴³

Combining these two strategies leads us to a handy model system that favours the dissociative pathway and can adapt its geometry through curving, as illustrated in the insets of Fig. 2(a). This system is denoted below as 2Co-N₄, emphasising its nature as an M-N-C dual-atom site catalyst. A key advantage of the 2Co-N₄ model is the linear dependence of energy on the squared curvature, i.e. inverse pore radius as shown in Fig. 2(a). In contrast, other geometry-adaptive catalysts, such as Pt nanoparticles, have a more complex energy-geometry relationship.44 As detailed below, linear dependence facilitates the modulation of individual adsorption

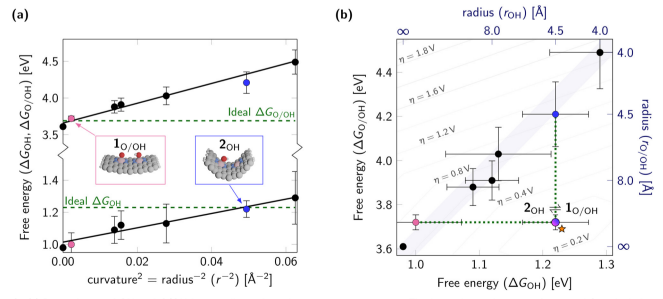


Fig. 2 (a) Dependence of OH and O/OH intermediate adsorption energy on curvature. The inset shows the states $\mathbf{1}_{\text{O/OH}}$ and $\mathbf{2}_{\text{OH}}$ depicting the adsorption of intermediates on the 2Co-N₄ model with selected curvature. (b) Top-view counter-map of the overpotential volcano for oxygen electrocatalysis. Grey lines represent overpotential contours. Coloured dots correspond to the states O/OH (pink - 21.2 Å), 20H (blue - 4.5 Å), and a geometry-adaptation between them (pink-blue gradient).

energies, thereby circumventing the scaling relations in a highly controlled manner.

Results

Adsorption free energies

The linear dependence in Fig. 2(a) gives rise to a new scaling relation for the dissociative mechanism: $\Delta G_{\text{O/OH}} = 2.93 \Delta G_{\text{OH}}$ + 0.70 eV,9 which constrains the predicted bifunctional overpotential to 0.62 V, surpassing the theoretical limit of 0.74 V for the associative mechanism.^{7,8}

For rigid catalysts, it can be generalised that any scaling relation results in a linear path on the corresponding activity volcano. Accordingly, each ΔG_{OH} - $\Delta G_{O/OH}$ point aligns within its uncertainty (see Model, methods, and FAIR data below) with the shaded line on the volcano's top view in Fig. 2(b). At a first glance, this linear scaling appears to corroborate numerous similar $\Delta G_{\rm OOH}$ vs. $\Delta G_{\rm OH}$ plots from existing literature, $^{2,6-8,10,45}$ where each point signifies a catalyst with a distinct chemical composition. At a second glance, the 2Co-N₄ model stands out. Unlike other scaling relations, in Fig. 2(b), all points share the same chemical composition of the 2Co-N₄ model. Still, the variable curvature produces a continuous range of $\Delta G_{\text{O/OH}} - \Delta G_{\text{OH}}$ points. Thus, the transition between points requires geometry adaptation of the 2Co-N₄ model, while in previous studies this would signify an alchemical transformation. This unique characteristic leads to a compelling question: "Can geometry adaptation between states of various curvature pave a nonlinear path to the apex of the activity volcano?" Answering this hypothetical question leads to the concept of geometry-adaptive electrocatalysis with a promise to bypass the scaling relations.

Geometry-adaptive electrocatalysis

Bypassing the scaling relations requires two states. For reaching the ideal catalyst these states correspond to the ideal adsorption energies of 1.23 and 3.69 eV for OH and O/ intermediates, respectively. In geometry-adaptive electrocatalysis on the 2Co-N₄ model, these two states are possible and marked as green dashed lines in Fig. 2(a). For verification, we selected states 1_{O/OH} and 2_{OH} in Fig. 2(a) that closely match the ideal adsorption energies. The resulting overpotential is below 0.1 V for the geometry adaptation between these states, which is visualised as an adjoining half-filled marker to the starred ideal catalyst in Fig. 2(b).

Such analysis focuses solely on thermodynamics and neglects barriers due to the reaction kinetics, surface coverage, reagents diffusion, the double layer structure, and ohmic resistance. Still, as the thermodynamics is the main limitation, it is inspiring to show that in principle, a thermodynamic overpotential of 0 V is achievable:

- · Because, unlike rigid catalysts, geometry-adaptive catalysts can bypass the scaling relations;
 - Specifically, through fine-tuning of the 2Co-N₄ model;
 - Generally, via analogous dynamic geometry-adaptation.

To draw an analogy from our findings, we have the following. While the quest for new rigid catalysts might be

compared to climbing straight paths on the activity volcano, geometry-adaptive electrocatalysis offers a bypass. It is like entering a tunnel at one state, moving directly to the central axis of the volcano, elevating to its apex, then returning, and finally emerging at another state. Fig. 2(b) illustrates the geometry-adaptive path with green dotted lines on the topview counter-map of the overpotential volcano for oxygen electrocatalysis. That path throughout the volcano is the essence of geometry-adaptive electrocatalysis.

The corresponding chemical cycle, including four combinations of intermediates and curvatures, is depicted in Fig. 3(a). This electrocatalytic cycle unites two reaction paths on the energy diagram in Fig. 3(b). Each path, on its own, suggests a modest catalyst, even though they incorporate states 1_{O/OH} and 2_{OH} with ideal adsorption energies. For instance, on the lower pink path, a significant kinetic barrier for the dissociation of O₂ hinders both OER and ORR.⁹ Yet, uniting these paths into a single cycle allows us to achieve almost ideal overpotential at the cost of curving the model.

In our DFT calculations, the 2Co-N₄ model is artificially curved under an assumption that, in reality, such geometry adaptation can be induced by external stimuli like electromagnetic radiation, temperature, electromagnetic fields, or pressure.46 With the employed DFT methods, we can not assess whether the substantial energy cost of 2.6 eV, required to curve the model from 21 Å to 4.5 Å, is reversible or includes additional overheads. Nevertheless, based on the results presented above and the discussion given below, we anticipate geometry adaptation to emerge as a new strategy for bypassing the scaling relations.

Pareto analysis

Reflections on the substantial curving energy prompted us to extend the analysis and consider the overpotential for all pairwise combinations of the curved models. The expanded analysis reveals a Pareto front, as depicted in Fig. 4(a), which trade-off visualises the between mechanical and geometry-adaptive electrochemical costs in oxygen electrocatalysis. Namely, for the 2Co-N₄ model, improving its catalytic activity in terms of overpotential comes at the expense of increasing curvature energy. The slope of the Pareto front is influenced by the energy required to curve the 2Co-N₄ model; as previously noted, the optimal overpotential is realised with a curvature energy of 2.6 eV. The Pareto analysis shows that, even for the 2Co-N4 model, there are promising candidates with reasonable curving energy and improved overpotential.

To put the Pareto front in perspective, let us turn the reader's attention to the approximate dependence of the overall efficiency of H2 production in an electrolyzer, followed by its oxidation in a fuel cell, on the bifunctional overpotential in Fig. 4(b). One can see that even a modest curving energy of 0.5 eV can reduce overpotential by 0.2 V, which translates to a near doubling of the energy efficiency compared to the state-of-the-art rigid catalysts (from ~25% to \sim 40%). Therefore, even modest geometry-adaptive catalysts

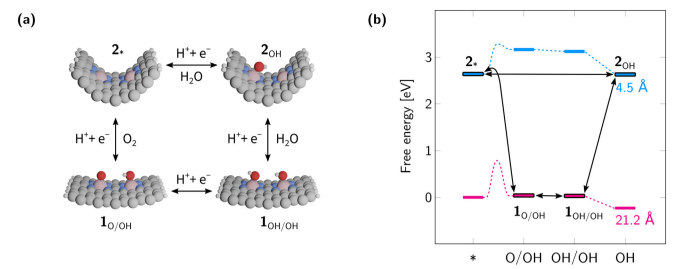


Fig. 3 (a) Chemical cycle for the geometry-adaptation between states 1_{O/OH}, 1_{OH/OH}, 2_{OH} and 2₊ for the O and OH intermediates adsorption at the 2Co-N₄ model of selected curvatures (1 and 2). Progressing clockwise leads to OER, whereas the counterclockwise direction results in ORR. (b) Free energy diagram at 1.23 V distinguishing between electrochemical steps (same colour) and mechanical steps (change in colour). Approximate barrier for dissociation step via proton-coupled electron transfer is shown schematically based on estimations for OOH dissociation.9

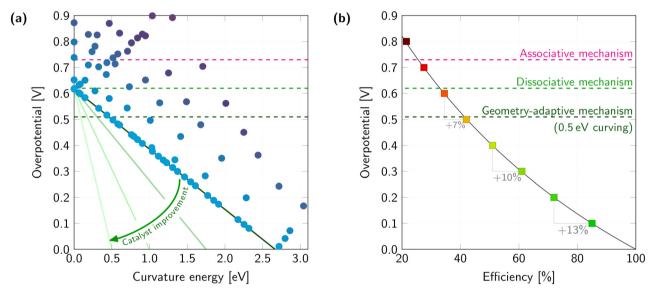


Fig. 4 (a) Pareto plot illustrating the trade-off between bifunctional overpotential ($\eta_{ORR} + \eta_{OER}$) and curvature energy. (b) Approximate dependence of overall efficiency (of H2 production in an electrolyser and its oxidation in a fuel cell) on the bifunctional overpotential

significantly boost the efficiency of oxygen electrocatalysis. This enhancement would benefit devices that rely on the OER and ORR, such as hydrogen electrolysers, fuel cells, and certain metal-ion batteries. As a result, these electrochemical devices could become more competitors to existing technologies, including internal combustion engines and traditional metal-ion batteries.

Discussion

Comparison of catalysts

In the diverse landscape of electrocatalysts, a systematic classification emerges on the basis of two criteria - structural

geometry adaptability and definiteness of the active sites. The former criterion divides catalysts into rigid and geometry-adaptive. As the name suggests, rigid catalysts retain their structure throughout electrocatalytic reactions. In contrast, geometry-adaptive catalysts tend to alter their geometry to optimise performance. The latter criterion categorises catalysts as either well-defined or stochastic. Welldefined catalysts possess active sites with precise atomic surroundings, whereas stochastic ones exhibit a degree of randomness in the atomic arrangement surrounding the active site.

Examples of well-known model systems corresponding to real catalysts are:

- Well-defined rigid platinum group metals and oxides are widely used in electrocatalysis due to their exceptional electrocatalytic activity controlled through geometry and composition. 47,48 Their structures, especially active sites, are well-defined and remain rigid during electrocatalytic reactions.
- Stochastic and rigid high-entropy alloys (HEAs) have a random atomic arrangement formed by mixing multiple principal elements.⁴⁹ Their stochastic nature means that the surrounding atoms of a given active site are random, providing a high degree of variability in electrocatalytic activity.
- Well-defined geometry-adaptive Fe-Cu dual-atom site in cytochrome c oxidase (CcO) follows the dissociative mechanism while reducing oxygen to water. 50,51 The difference in Fe-Cu distance in reduced and oxidised forms of the enzyme can be viewed as geometry adaptation.
- Stochastic geometry-adaptive M-N-C catalysts with dualatom sites, 52 exemplified by the 2Co-N₄ model (in this work), uniquely combine design with inherent randomness in electrocatalysis. Their intrinsic curvature can facilitate geometry adaptation. This adaptability and random atom site distribution define their distinct catalytic character.

Together, these catalysts represent a spectrum of properties, from the randomness of HEAs to the precision of enzymes, each offering unique advantages and challenges in electrocatalysis. M-N-C materials with 2Co-N₄ dual-atom sites emerge as a unique blend in this classification. These materials are inherently curved, relatively flexible, and have a stochastic distribution of catalytic sites.⁵³⁻⁵⁶ Each dual-atom site can potentially adopt geometry during reaction independently of others under external stimuli.46 However, mastering such geometry adaptation control (like in enzyme cofactors) is an open challenge. The presented 2Co-N₄ model is just a first step in unravelling geometry-adaptive mechanisms in artificial catalysts.

Perspective on circumventing the scaling relations

Let us put the challenge of implementing geometry-adaptive electrocatalysis into a broader context of circumventing scaling relations. There are five general strategies:

- 1. Tuning by varying the OH adsorption energy;^{57–59}
- 2. Breaking by stabilising OOH relative to OH; 1,7,60
- 3. Switching from the associative to the dissociative mechanism;61-65
 - 4. Pushing by destabilising O relative to OH;^{9,61}
- 5. Bypassing by completely decoupling the adsorption energies.

While tuning and breaking the OH-OOH scaling relation are well-established in current state-of-the-art, 1,7,34,57-60,66 switching mechanisms represent the cutting edge of research, with confirmed experiments. 62,64,65 knowledge, pushing the OH-O/OH scaling relation is computationally conceptualised, 9,61 and awaits experimental verification in direct relation to research on switching. In this work, we propose the geometry adaptation concept as a candidate for the bypassing strategy.

The rhetorical question "How can this adaptation occur at a frequency matching catalytic turnover?" inspires us to look closer at the CcO enzyme. Regarding the turnover frequency, CcO surpasses all known artificial catalysts by orders of magnitude.³⁸ The active site of CcO favours the dissociative mechanism and undergoes structural changes during catalysis. 67-70 Therefore, we speculate that CcO follows geometry-adaptive catalysis and bypasses scaling relations. Due to the trade-off between costs for geometry adaptation and overpotential, CcO is not the ideal catalyst, yet it shows a record low compromise overpotential of 0.19 V.71

biocatalysis, including Although CcO switching configurations and pumping protons, is a marvel of complexity, it also highlights the limitations inherent in biological systems. Unlike biocatalysis, electrocatalysis is not bound by the constraints of biological evolution and can utilise a broader range of chemical elements and create more diverse and exotic structures; it can also apply effects that are rare or non-existent in biological systems, such as electromagnetic fields, extreme pressures or temperatures. Moreover, human engineering can often surpass natural processes in terms of efficiency, scalability, and specificity.

For inspiration, consider the ferroelectric model, which demonstrates high-frequency switching between states during reaction.⁷² Besides, alternative models and mechanisms for geometry adaptation could sustain a higher frequency. For example, using rotational degrees of freedom in nanotubes and buckyballs (see Fig. 5) offers a less energy-intensive means of adjusting the active site geometry. addition, controlling geometry using cis-trans photoisomerisation could induce localised curving around active sites in M-N-C-like materials.⁷³ The possibilities are only limited by the laws of nature and human creativity. As a source of ideas, we recommend a recent review by Wang et al. with examples of geometry adaptations in electrocatalysts, highlighting many opportunities for further advancements in this field.74

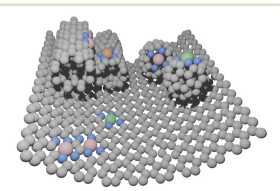


Fig. 5 Visualisation of M-N-C single- and dual-atom site models within a graphene layer, nanotubes, and buckyballs. The dissociative oxygen electrocatalysis on these sites can be controlled by curving and rotating.

Further considerations

Communication

This study builds upon the emphasis on the geometric effect in electrocatalysis, recently recognised as an independent and significant factor.⁷⁵ On the one hand, models focusing on the geometric effect for single-atom site catalysts (in particular with variable curvature) help reach the apex of the "associative mechanism" activity volcano in Fig. 1(a). One of the pioneering works, by Xie et al. in 2018,76 demonstrated an energy-curvature relationship (like in Fig. 2) employing a model similar to 2Co-N₄. More recent works on M-N-C- and metal phthalocyanine-decorated carbon nanotubes, as well as curved graphene, have predicted enhanced performance attributed to the curvature-induced strain effect.77-79 The strain effect was known for a long time and nowadays is used in modelling catalysts for oxygen evolution and reduction reactions, 80,81 as it alters the OH-OOH scaling relation. 82

On the other hand, models of curved in-pore, slit-pore, biphenylene, difullerenes, dimicrocycles, and diporphyrins catalysts have proposed for dual-atom site been catalysts. 9,10,39,61,83,84 All referred models, firstly, favour the dissociative mechanism and, secondly, can adjust geometry for specific intermediates. Therefore, such models can be candidates for geometry-adaptive electrocatalysis.

Notably, Wan et al. overviewed the fundamentals of electronic and geometric effects by comparing metal with dual-atom site catalysts to conclude: "the direct way to achieve this (higher intrinsic catalytic activity) is to design catalysts that go through a direct dissociative mechanism, bypassing the OOH intermediate". 10 Also highlighted are the main complication beyond thermodynamic analysis - the kinetic barriers. 85,86 Specifically, the dissociative mechanism has a notable kinetic barrier for the O2 reduction via a proton-coupled electron transfer to O and intermediates.⁸⁷ Using the Brønsted-Evans-Polanyi relation and data from the literature, 9,10 we approximated the kinetic barrier as schematically shown in Fig. 3(b). Due to the nearoptimum inter-site distance in the state 2, the barrier for both O2 and OOH dissociation is low. In particular, the geometry-adaptive mechanism in the 2Co-N4 model is adjustable by curving. To fully understand the kinetic aspects of proton-coupled electron transfer during dissociation, DFTbased molecular dynamics is essential, 87,88 which, while beyond the scope of this study, would further reinforce the concept of geometry-adaptive electrocatalysis.

Conclusions

We have explored an approach to elevate the efficiency of oxygen electrocatalysis using a geometry-adaptive model system and density functional theory methods. We show that geometry adaptation helps to bypass the scaling relations by adjusting the geometry of the model system and adapting the geometry during the catalytic reaction.

The introduced model system is a dual-atom site 2Co-N₄ with a unique geometric effect - a linear dependence

between geometry and the adsorption energy of reaction intermediates. In the example of the 2Co-N₄ model, we highlight the difference between novel geometry-adaptive and classical rigid catalysts. Namely, the geometry adaptation during reactions can bypass the fundamental limitation due to scaling relations.

Our computational modelling of the 2Co-N₄ model revealed several key insights:

- The geometric effect allows reaching ideal adsorption energies for reaction intermediates.
- Geometry adaption during oxygen evolution/reduction reactions allows to reach the ideal potential for oxygen electrocatalysis.
- The direction for improving geometry-adaptive catalysts towards optimal catalysts is given by the Pareto front - a trade-off between costs for geometry adaptation and overpotential (deviation from the thermodynamic potential).

With these insights, we hope to inspire further research on geometry-adaptive catalysts to discover the true ideal oxygen electrocatalysts - elevating to the apex of the activity volcano - and address the current lack of significant progress in the experimentally observed activities.

Model, methods, and FAIR data

The 2Co-N₄ model is an M-N-C dual-atom site catalyst of a constant composition and variable curvature. Two distinct geometry states of the model, nearly planar and curved, are depicted in Fig. 2(a).

DFT calculations for intermediates adsorbed on the 2Co-N₄ model followed standard practices, as referenced in previous works. 61,89 Spin-polarised DFT calculations were performed with the GPAW 22.8 and ASE 3.22.1 software using the revised Perdew-Burke-Ernzerhof (RPBE) functional. 90-92 For optimisation and energy calculations, $3 \times 1 \times 1$ k-point sampling, grid spacing of 0.18 Å was used. After curving, a 5 Å vacuum layer was added in all non-periodic directions. Only edge carbon atoms were constrained to define the curvature, while all other atoms, including intermediates, were fully optimised below 0.1 eV Å⁻¹. The dissociative mechanism was the focus of this study:

$$2* + O_2 + (H^+ + e^-) \rightarrow *O + *OH$$
 (1)

$$*O + *OH + (H^{+} + e^{-}) \rightarrow *OH + *OH$$
 (2)

$$*O + * + (H^{+} + e^{-}) \rightarrow *OH + *$$
 (3)

*OH + * + (H⁺ + e⁻)
$$\rightarrow$$
 2* + H₂O (4)

where * indicates one of two adsorption sites on the 2Co-Nx model of variable curvature.

The free energy (G) of each intermediate was calculated as:

$$G = E + ZPE + TS + S \tag{5}$$

where E is the DFT energy with the D4 correction, ZPE is the zero-point energy, TS is the entropy correction, S is the solvation correction. 8,61,93 The uncertainty for the adsorption energies in Fig. 2 are estimated from the variance of the BEE ensemble predictions, 94,95 as implemented in ASE. 91 The adsorption free energies (ΔG) of each intermediate were obtained for the dissociative mechanism as:

$$\Delta G_{\rm O/OH} = G_{\rm O/OH} - G_* + \frac{3}{2}G_{\rm H_2} - 2G_{\rm H_2O} - 3eU \tag{6}$$

$$\Delta G_{\text{OH/OH}} = G_{\text{OH/OH}} - G_* + G_{\text{H}_2} - 2G_{\text{H}_2\text{O}} - 2eU$$
 (7)

$$\Delta G_{\rm OH} = G_{\rm OH} - G_* + \frac{1}{2} G_{\rm H_2} - G_{\rm H_2O} - eU \tag{8}$$

where the computational hydrogen electrode was used to calculate the energy of $(H^+ + e^-)$ at potential U as $\frac{1}{2}G_{H_2} - eU$ with e being the elementary charge.⁵

The OER and ORR overpotential values were evaluated in terms of energy differences between reaction stages in Fig. 3:

$$\eta_{\text{ORR}} = 1.23 \text{ V} - \min(\Delta G_{1-4})/e, \, \eta_{\text{OER}} = \max(\Delta G_{1-4})/e - 1.23 \text{ V}$$
 (9)

These expressions define the volcano in Fig. 1(a).^{7,61}

FAIR data on the modelled structures, total energy values, and analysis scripts are accessible on the webpages https:// chem.ku.dk/research_sections/nanochem/theoreticalelectrocatalysis/ and https://nano.ku.dk/english/research/ theoretical-electrocatalysis/katladb/geometry-adaptive-catalysis/.

Author contributions

R. Cepitis: conceptualisation, data curation, formal analysis, investigation, methodology, writing - original draft; V. Ivaništšev: conceptualisation, software, supervision, validation, visualisation, writing - review & editing; J. conceptualisation, validation, Rossmeisl: resources, visualisation; N. Kongi: supervision, funding acquisition, writing - review & editing, project administration.

Conflicts of interest

There are no conflicts to declare.

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