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Research progress on layered metal oxide electrocatalysts for an efficient oxygen evolution reaction

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Hydrogen, highly valued for its pristine cleanliness and remarkable efficiency as an emerging energy source, is anticipated to ascend to a preeminent status within the forthcoming energy landscape. Electrocatalytic water splitting is considered a pivotal, eco-friendly, and sustainable strategy for hydrogen production. The substantial energy consumption stemming from oxygen evolution side reactions significantly impedes the commercial viability of water electrolysis. Consequently, the pursuit of a cost-effective and efficacious oxygen evolution reaction (OER) catalyst stands as an imperative strategy for realizing hydrogen production via water electrolysis. Layered metal oxides, owing to their robust anisotropic properties, versatile adjustability, and extensive surface area, have emerged as suitable candidates for OER catalysts. However, owing to the distinctive attributes of layered metal oxides, ongoing investigations into these materials are slightly fragmented, lacking universal consensus. This article comprehensively surveys the recent advancements in layered metal oxide-based OER catalysts, categorized into single metal oxides, alkali cobalt oxides, perovskites, and miscellaneous metal oxides. Initially, the main OER intermediate reaction steps of layered metal oxides are scrutinized. Subsequently, the design, mechanism, and application of several pivotal layered metal oxides in the OER are systematically delineated. Finally, a summary is provided, alongside the proposal of future research trajectories and challenges encountered by layered metal oxides, with the aspiration that this paper may serve as a valuable reference for scholars in the field.

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# 1. Introduction

As technological advancements burgeon and societal evolution unfolds, the echelons of global industrialization ascend ever higher. The trajectory of industrial advancement remains inexorably intertwined with humanity's exploitation and utilization of the natural world, leading to increasingly prominent damage to the natural environment. Presently, the main energy consumption stems predominantly from non-renewable resources, including coal mines, oil, and natural gas, notwithstanding the growing integration of eco-friendly alternatives such as wind, solar, and tidal energies.<sup>1-4</sup> Nevertheless, they are often limited by external factors such as natural climate and cannot be utilized for a long time, and their contribution to the overall energy framework remains relatively minimal.<sup>5,6</sup> Therefore, there emerges an imperative need to develop a novel sustainable green energy, supplanting fossil fuels, thus protecting the earth's environment. Hydrogen energy, as a new type of green energy, has a wide range of sources, high heat of combustion value, clean and pollution-free nature, and an intrinsic capacity for energy storage, which enable it to play a pivotal role in the prospective energy landscape.<sup>7,8</sup> Currently, two primary avenues exist for hydrogen production: one involves harnessing fossil fuels to generate hydrogen, such as gas reforming, oil refining, coal gasification, and alcohol cracking, whereas the other entails green hydrogen synthesis *via* the electrolysis of water using electricity or photoinduced decomposition.<sup>9–14</sup> Conversely, electrolytic water splitting for hydrogen generation stands out for its simplicity, diversity in electrical energy sources, and myriad benefits such as heightened product purity and environmental friendliness, thus positioning itself as a promising avenue for hydrogen production.<sup>15–17</sup>

In the process of water electrolysis, two pivotal half-reactions unfold at the anode and cathode: the two-electron hydrogen evolution reaction (HER) and four-electron oxygen evolution reaction (OER).<sup>18,19</sup> Utilizing alkaline electrolytes as a case study,<sup>20</sup> the reaction equations are as follows:

$$\begin{split} & \text{HER}: 2\text{H}_2\text{O}(\text{l}) + 2\text{e}^- \to \text{H}_2(\text{g}) + 2\text{OH}^-(\text{aq}) \\ & E^{\bigcirc}_{\text{H}_2} = 0.00\,\text{V}(\nu s.\,\text{RHE}) \end{split} \tag{1}$$

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$$\begin{array}{l} {\rm OER}: 4{\rm OH}^-({\rm aq}) \to 2{\rm H}_2{\rm O}({\rm l}) + {\rm O}_2({\rm g}) + 4{\rm e}^- \\ E_{{\rm O}_2}^\circ = 1.23\,{\rm V}(\nu s.\,{\rm RHE}) \end{array} \tag{2}$$

$$\begin{aligned} & \text{Overall}: 2\text{H}_2\text{O}(l) \rightarrow \text{O}_2(g) + 2\text{H}_2(g) \\ & E^{\bigcirc}_{\text{Overall}} = 1.23 \, \text{V}(\text{redox}). \end{aligned}$$

Theoretically, water decomposes into H<sub>2</sub> and O<sub>2</sub> and requires a voltage of 1.23 V between the cathode and anode to facilitate the transfer of 4e<sup>-.21</sup> However, owing to the intricate nature of intermediate reactions, the conversion process of intermediary products must surmount certain potential barriers to propel the progression of both cathodic and anodic reactions.<sup>22</sup> To mitigate the Gibbs free energy of the reaction process in kinetics, thereby alleviating the overpotential of the reaction, the utilization of catalysts has surfaced. Given the involvement of four-electron transfers in the OER process, it is widely acknowledged that OER proceeds at a significantly slower pace than that of the HER in kinetics.<sup>23,24</sup> Hence, the development of efficient OER catalysts stands as pivotal in enhancing the overall water decomposition efficiency. In the nascent stages of research, noble metal-based catalysts attracted extensive attention due to their remarkable activity and stability. Notably, IrO2 and RuO2 have found widespread application in OER processes and serve as primary commercial catalytic materials.<sup>25,26</sup> Consequently, researchers are developing novel OER catalysts from three perspectives: (1) to reduce dependence on precious metals, (2) to develop precious metalbased compounds, and (3) to develop non-precious metalbased materials.<sup>27-29</sup> Since the electrolysis of water for hydrogen production has entered the field of research, a large number of metal-based OER catalysts have been developed, spanning metal oxides, sulfur compounds, nitrides, phosphides, and beyond.<sup>30-33</sup> Two-dimensional materials including metal halides, metal oxides, metal hydroxides have attracted much attention due to their significant advantages in exposing multiple active sites.<sup>34–36</sup>

This article focuses predominantly on summary OER catalysts grounded in layered metal oxides (LMOs). Layered metal oxides represent a distinctive category of solid materials characterized by pronounced anisotropy across their basal and edge planes, thereby conferring upon them unexpected chemical and physical attributes.<sup>37,38</sup> Moreover, LMOs present a capacity for undergoing low-temperature chemical transformations devoid of compromising the integrity of covalent bonds within the layers.<sup>39</sup> These metal oxides may assume a morphology comprising a few or single layers crafted through specific methodologies, or they may show as stacks of electrically neutral metal oxide layers. However, more commonly, they coexist with charged layers interwoven with cations or anions, or their metal oxide sheets alternate with the middle layer of covalent bonds.40-43 LMOs boast several advantageous traits: (i) thermodynamic stability and difficulty in breaking covalent bonds within the layer.44,45 (ii) In most cases, in aqueous and highly oxidizing environments, metal oxides readily transition into corresponding metal (oxygen) hydroxides, which serve as genuine OER active species.<sup>46</sup> (iii) LMOs exhibit compositional flexibility and adaptable intercalation groups, thereby engendering a diverse coordination environment at their active sites, thereby affording excellent OER activity and stability.<sup>40,47,48</sup>

In this comprehensive review, we scrutinize the recent strides in researching the materials and structures of LMOs as OER catalysts. Initially, we delineate three distinct evolutionary mechanisms operative in the OER process. Subsequently, we provide a concise overview of several conventional and newly discovered layered metal oxide OER catalysts, encompassing single metal oxides, alkali cobalt oxides, perovskites,  $V_2O_5 \cdot nH_2O_1$ , cobalt phyllosilicate, brownmillerite, and weberite. On this basis, we expound upon methodologies for modifying layered metal oxides, elucidate key factors contributing to enhancing their activity or stability, and point out the gaps that still exist in research. Finally, we summarize the relationship between the structures of layered metal oxides, the triggering of OER mechanisms, and the factors enhancing electrocatalytic performance. Moreover, we outline the future research directions and challenges confronting layered metal oxides (Fig. 1). This endeavor aims to furnish valuable insights into the application of layered metal oxides in the OER domain, thereby accelerating the commercialization of hydrogen production through water electrolysis.



**Fig. 1** Schematic diagram of the oxygen evolution reaction process of LMOs from basic materials and reaction mechanisms to OER performance. (AEM: adsorbate evolution mechanism; LOM: lattice oxygen evolution mechanism; OPM: oxide path mechanism.) Adapted with permission from ref. 73. Copyright 2020, the Royal Society of Chemistry. Adapted with permission from ref. 66. Copyright 2021, Elsevier. Adapted with permission from ref. 93. Copyright 2022, American Chemical Society. Adapted with permission from ref. 106. Copyright 2017, WILEY. Adapted with permission from ref. 106. Copyright 2022, American Chemical Society. Adapted with permission from ref. 107. Copyright 2023, Springer Nature.

### 2. Hydrogen economy

With rapid development of global industries, nations are strategically elevating hydrogen energy to a paramount position on their agendas. The United States was the first to propose the concept of using hydrogen energy as a basic energy carrier to build a future energy society. As early as 2001, the US Department of Energy issued the "2030 Vision for America's Transition to a Hydrogen Economy", which pointed out that hydrogen will replace fossil fuels as the main energy source supporting the global economy in the future, forming a market operation system based on the production, storage, transportation, and application of hydrogen energy, known as the "Hydrogen Economy".<sup>49</sup> Moreover, numerous nations including Germany and China have crafted extensive blueprints for the long-term utilization of hydrogen energy technology, positioning it strategically within their agendas. Electrolytic hydrogen production technology, leveraging renewable green energy sources, stands poised to effectively mitigate intermittent energy consumption challenges posed by solar and wind energy. This technology assumes a pivotal role in the storage, conversion, and optimal utilization of clean energy resources.<sup>50</sup> Nonetheless, the issue of hydrogen persistent production costs remains a central issue thwarting the widespread commercialization of electrolytic water hydrogen production technology. In terms of green hydrogen production capacity, the European Union aims to establish an electrolysis system of 40-60 GW by 2030. The relevant strategy also points out the specific goal of achieving a hydrogen production capacity of 1 million  $t \cdot a^{-1}$  by 2030. The Netherlands also pointed out that, based on the existing 10 MW electrolytic cell system, a 3-4 GW green hydrogen production plant will be further constructed before 2030.51,52 Nevertheless, to realize sustained production of high-quality and low-cost hydrogen, resolving the issue of energy consumption in the anode's OER process is imperative. Therefore, designing and fabricating highly active and enduringly stable catalysts emerge as pressing imperatives in attaining the envisioned hydrogen energy landscape.

# 3. OER mechanisms for transition metal oxide electrocatalysts

Mastering the relationship between catalyst performance and reaction mechanism is crucial for guiding catalyst design. Currently, three distinguished evolutionary mechanisms for OER catalytic processes have been acknowledged.53-55 The most common reaction mechanisms are as follows: (i) adsorbate evolution mechanism (AEM), (ii) lattice oxygen-evolution mechanism (LOM), and (iii) the novel mechanism discovered in recent years: oxide path mechanism (OPM). A fundamental relationship exists between the catalytic reaction mechanism and the catalytic active site. Moreover, various electrolytes induce disparities in the initial reactants, thereby exerting a discernible influence on the reaction process. In alkaline solutions, a substantial concentration of ionized OH<sup>-</sup> facilitates the onset of the OER process. Within this context, a concise overview of the reaction process alongside illustrative examples is presented to elucidate the diverse mechanisms operative in alkaline electrolyte environments.

#### 3.1 Adsorbate evolution mechanism (AEM)

As shown in Fig. 2a, during the OER process, the AEM pathway involves four electron-proton transfer steps occurring at the metal-ion centers exposed to the catalyst surface.<sup>56</sup> Initially, the metal active site combines with the OH<sup>-</sup> ion in the electrolytes and evolves into \*OH attached to the metal (where \* denotes the metal site). Subsequently, the intermediate \*OH reacts with OH<sup>-</sup> in the electrolyte to form an \*O intermediate *via* deprotonation. Following this, an another OH<sup>-</sup> ion from electrolytes combines with the \*O intermediate to form the intermediate \*OOH. Further deprotonation evolves into O<sub>2</sub> molecules, which then escape from the metal site, leaving a vacancy that is then replenished with OH<sup>-</sup> in the electrolyte solution.<sup>57</sup> Within this cycle, the OER process catalyzed by the AEM exhibits a circular nature, yielding the intermediates \*OH, \*O, and \*OOH.

Ideally, the catalytic process following AEM should have an equal Gibbs free energy at each reaction step, totaling 1.23

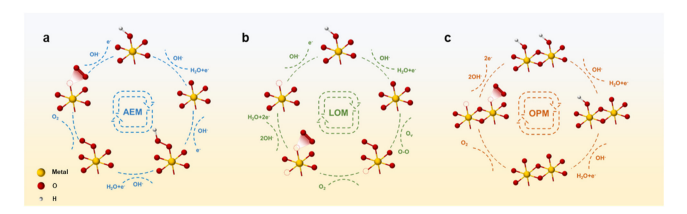


Fig. 2 Schematic diagram of (a) AEM, (b) LOM, and (c) OPM in alkaline OER process.

eV.58 However, a fixed scaling relationship exists between the binding energies resulting from \*OH deprotonation and \*OOH formation. Therefore, there exists an almost constant energy difference  $\Delta G_{*OOH} - \Delta G_{*OH} = 3.2$  eV. Consequently, a deviation arises between the actual and theoretical free energies, resulting in a minimum overpotential range of 0.2-0.4 V for the AEM process.<sup>59,60</sup> The variance in Gibbs free energy at each step leads to the presence of a rate-determining step (RDS) within the AEM process, typically associated with either \*OH deprotonation or \*OOH formation, thereby influencing the oxygen evolution reaction (OER) potential.<sup>61</sup> Meanwhile, owing to the constant energy difference of  $\Delta G_{*OOH} - \Delta G_{*OH} = 3.2$  eV, the OER overpotential is described by the  $(\Delta G_{*O} - \Delta G_{*OH})$ value as the OER activity descriptor. Typically, when the  $(\Delta G_{*O} - \Delta G_{*OH})$  value is around 1.6 eV, it indicates an optimal oxygen binding strength conducive to optimal OER activity. Below this value, the binding strength of oxygen will be too strong, hindering the subsequent formation of \*OOH, whereas surpassing this threshold renders oxygen binding strength too weak, impeding \*OH deprotonation.<sup>62</sup>

Subsequently, Hammer and Nørskov proposed the d-band theory based on first-principles calculations, utilizing the d-band center as a single parameter to evaluate the effect of clean metal surfaces on the catalytic activity of metal groups.<sup>63</sup> Based on their research, the d-band center exhibits a close correlation with the stability of surface atoms/molecules and the transition state energy of surface processes. The shift of the d-band center of the active site transition metal towards the Fermi level serves to enhance the adsorption of intermediates. Conversely, a greater distance of the d-band center from the Fermi level results in the weakening of the adsorption of intermediates.<sup>64,65</sup> Inspired by the d-band center theory, Dai et al. discovered that the intrinsic activity of layered alkali metal oxides can be regulated by adjusting the CoO<sub>6</sub> octahedron, which is shared by the inner edge of the layer and spacing/strain between CoO<sub>2</sub> layers. The authors investigated the modulation of the OER activity of Na<sub>0.7</sub>CoO<sub>2</sub> layered oxides by Ag, Cu, and Ce doping and cation removal. The results indicate that the lattice strain introduced by the cation removal process leads to a positive shift in the d-band center of the surface Co atom, thereby modulating  $\Delta G_{*OH}$  and facilitating the deprotonation process towards a moderate free energy, consequently lowering the energy barrier during the reaction process.66

#### 3.2 Lattice oxygen-evolution mechanism (LOM)

Through extensive investigation into the OER mechanism, researchers have uncovered that oxygen atoms constituting oxygen molecules may arise not only from water molecules but also from the lattice oxygen evolution of metal oxides (Fig. 2b).<sup>67</sup> The electronic configuration of the active sites in such OER catalysts typically deviates from the d-band center state conducive to maintaining Gibbs free energy equilibrium in the AEM process. The sequential stages of the OER process diverge from those observed in the AEM. Initially, OH<sup>-</sup> in the electrolyte adsorbs on metal active sites, evolving into \*OH. Go

through the deprotonation step to form the intermediate \*O. In contrast to the continuation of OH<sup>-</sup> adsorption to generate the \*OOH intermediate in the AEM process, \*O binds to lattice oxygen within the catalyst body, liberating it as O<sub>2</sub>. Due to the deficiency of lattice oxygen, oxygen vacancies are generated within the principal lattice structure of the catalyst.<sup>57</sup> Subsequently, two distinct methods are employed to replenish oxygen vacancies. On the one hand, other oxygen atoms in the catalyst body diffuse to occupy vacancies, and on the other hand, defects are filled through OH<sup>-</sup> adsorption and deprotonation in the electrolyte. This particular OER mechanism is referred to as LOM due to its involvement of oxygen atoms within the catalyst body in addition to those present in the electrolyte solution. In addition, the LOM process bypasses the limitation of intermediate \*OOH formation, thus, layered oxides based on LOM hold promise for achieving enhanced OER activity.<sup>68</sup> However, attention to the pervasive issue of heavy metal leaching, commonly associated with LOM, proves challenging.<sup>69</sup> For example, Ge et al. synthesized an excellent OER catalyst by loading single-atom Ir sites onto layered MnO<sub>2</sub>. More precisely, the introduction of single Ir atoms triggers the activation of lattice oxygen. Compared to the bulkphase IrO<sub>2</sub>, the Ir-O bond length of the Ir single atom dispersed on the surface of MnO<sub>2</sub> is shortened, which is more suitable for the chelation structure. Authors directly demonstrated that the LOM process on the isolated Ir site was opened through in situ methods. Therefore, it has better OER activity, which is more than 42 times that of industrial IrO<sub>2</sub>. Furthermore, the activation of local lattice oxygen mitigates significant ion migration during the OER process, thereby preventing catalyst structure degradation. As a result, the catalyst has a stability up to 650 hours.<sup>70</sup>

#### 3.3 Oxide path mechanism (OPM)

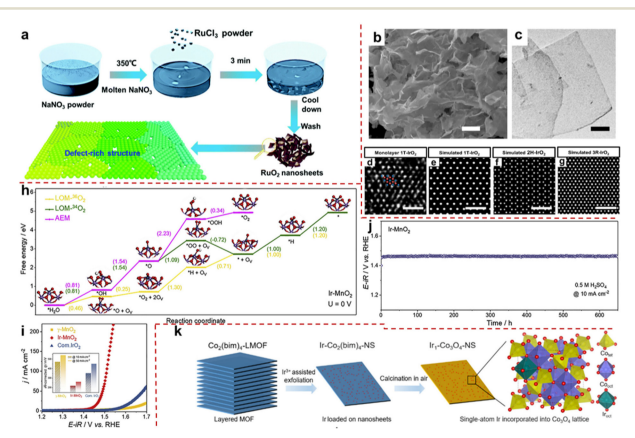
Apart from the AEM and LOM processes, there exists the OPM characterized by OER activity and stability surpassing theoretical thresholds.<sup>40,71</sup> The OPM pathway is different from other reaction processes in that one unit of the reaction process occurs simultaneously at two metal active sites, facilitating the direct coupling of O-O radicals. In the OER process driven by the OPM, only intermediates \*O and \*OH are generated, devoid of oxygen vacancy defects and reaction intermediates \*OOH (Fig. 2c).<sup>53,55</sup> The OPM process needs to work collaboratively through active metal sites at appropriate positions to deprotonate and trigger the coupling of \*O and other \*O radicals to generate O2. Therefore, stringent demand exists for the geometric arrangements of metal active sites. For example, Shigeo Mori et al. introduced a novel structure of perovskite  $CaCu_3Fe_4O_{12}$  (CCFO). Compared to the traditional ABO<sub>3</sub>-type perovskite, this structure not only stabilizes via covalent Fe-O-Cu bonds but also diverges from the conventional AEM pathway, embracing the OPM with bimetallic active sites. During the OER process, effective electron transfer to Fe ions occurs, facilitated by robust Fe-O covalent bonds, with the reaction proceeding via the oxidation-reduction of Fe<sup>4+</sup>/Fe<sup>5+</sup> ions.<sup>72</sup> Moreover, owing to the unique perovskite structure, the

Fe–Fe atomic distance remains relatively short, naturally facilitating the operation of the OPM.

# 4. Layered oxides for OER catalysts

#### 4.1 Single metal oxides

Currently, two primary approaches exist for the modification of layered single metal oxide OER catalysts: (i) morphological regulation. Through specialized methods and conditions, conventional single metal oxides are transformed into twodimensional layered structures, primarily aimed at enhancing surface exposure, augmenting active sites, and consequently, fostering heightened OER activity. (ii) Leveraging layered oxides as carriers facilitates the facile dispersion of precious metal units onto their surfaces, thereby diminishing the precious metal loading while ameliorating the electronic structure of metal sites, thus favoring the realization of excellent OER activity. Li *et al.* prepared ultra-thin single-layer or few-layer  $RuO_2$  nanosheets ( $RuO_2$  NSs) by a simple molten salt method using NaNO<sub>3</sub> as a template at specific temperatures (Fig. 3a). The  $RuO_2$  NS surface has abundant Ru vacancies, significantly weakening the binding energy of \*O, thereby reducing the energy barrier of the conversion process from \*O to \*OOH and improving OER performance. At a current density of 10 mA cm<sup>-2</sup>, the overpotential is extremely low at 199 mV. The excellent performance is attributed to the exposure of high surface area in layered structures and the influence of defect introduction on the electronic structure of active centers.<sup>73</sup> Shao et al. reported the synthesis of pure-phase IrO<sub>2</sub>: 1T-phase IrO<sub>2</sub> (1T-IrO<sub>2</sub>) achieved through a combination of mechanochemistry and heat treatment in a strongly alkaline medium. As shown in Fig. 3b and c, the SEM and TEM images reveal that the synthesized IrO<sub>2</sub> possesses an ultra-thin layered structure. In addition, its atomic arrangement aligns with the crystal structure of 1T phase (Fig. 3d). IrO<sub>2</sub> materials usually have three phase structures: 1T, 2H, and 3R. Unlike the two closely packed stacking arrangements of 2H and 3R, the 1T phase exhibits a distinctive layered structure characterized by AA stacking (Fig. 3e-g). Compared to  $IrO_2$  in the rutile phase, the special phase structure of 1T-IrO<sub>2</sub> facilitates the generation of \*OH species during the OER process, consequently enhancing the OER activity. At a constant current density of 10 mA cm<sup>-2</sup>, the overpotential is as low as 197 mV.



**Fig. 3** (a) Schematic route of the synthesis of the defect rich monolayer  $RuO_2$ . Adapted with permission from ref. 73. Copyright 2020, the Royal Society of Chemistry. (b and c) SEM and TEM images of 1T-Ir $O_2$ . (d) HAADF-STEM image of single-layer 1T-Ir $O_2$  atomic arrangement. Ir and O are represented by blue and red spheres. (e–g) Simulated TEM images of  $IrO_2$  atomic arrangement in 17, 2H, and 3R phases. Adapted with permission from ref. 74. Copyright 2020, Springer Nature. (h) Gibbs free energy calculation of the OER process shows Ir-MnO<sub>2</sub> (100) at 0 V. (i) LSV curve of Ir-MnO<sub>2</sub> in 0.5 M H<sub>2</sub>SO<sub>4</sub>. The illustration shows the overpotential values at different current densities. (j) Stability curve of Ir-MnO<sub>2</sub> at a constant current density of 10 mA cm<sup>-2</sup>. Adapted with permission from ref. 70. Copyright 2021, Cell Press. (k) Schematic route and structure of Ir-Co<sub>3</sub>O<sub>4</sub> layered nanosheet synthesis. Adapted with permission from ref. 77. Copyright 2023, American Chemical Society.

Assembled in a proton exchange membrane device, it exhibits excellent stability for 126 hours at a high current density of 250 mA cm<sup>-2</sup>.<sup>74</sup>

Typically, the utilization of layered pure-phase single metal oxides in OERs is confined to precious metal oxides, which suffer from the disadvantages of scarce resources, exorbitant costs and notable constraints. Therefore, there exists a pressing need to pioneer the development of non-noble metal layered oxides for their utilization in the field of OER catalysis. Given the intrinsic activity constraints of non-precious metal oxides, it remains imperative to integrate them with minute quantities of noble metals to bolster performance.75,76 For example, ordinary IrO2 follows the AEM process, which is slow and requires overcoming large energy barriers. The IrMnO<sub>x</sub> obtained by doping transition metal Mn into IrO<sub>2</sub> can trigger the LOM process, avoid the formation of the intermediate \*OOH and greatly reduce reaction energy consumption. However, the lattice oxygen deficiency caused by its OER process is irreversible and cannot serve stably for a long time. Therefore, the development of high-activity and long-term stability OER catalysts has become the main research direction. Ge et al. achieved the activation of lattice oxygen in Ir unit point catalyst (denoted as Ir-MnO<sub>2</sub>) for the first time. Compared with bulk IrO<sub>2</sub>, the dispersed Ir sites of atoms in acid-stabilized  $\gamma$ -MnO<sub>2</sub> adapt to the chelating structure of the latter. The author demonstrated through in situ <sup>18</sup>O isotopelabeled differential electrochemical mass spectrometry that the LOM on the separated Ir site was opened, which was due to a shortened Ir-O bond length (5% contraction compared to IrO<sub>2</sub>). Density functional theory calculations indicate that the Ir-MnO<sub>2</sub> triggered the LOM process, and the filling of lattice oxygen, whether from within the crystal or electrolyte solution, results in significantly lower RDS Gibbs free energy (1.09 eV, 1.3 eV) in the OER process compared to the AEM process (2.23 eV) (Fig. 3h). Therefore, Ir-MnO<sub>2</sub> has a good low overpotential (218 mV (a) 10 mA cm<sup>-2</sup> at 0.5 M H<sub>2</sub>SO<sub>4</sub>) (Fig. 3i). In addition, due to the activation of local lattice oxygen, the migration of a large amount of ions is inhibited during the OER process, which hinders the collapse of the catalyst structure. Therefore, after 650 hours of durability testing, the catalyst can still maintain its activity well (Fig. 3j).<sup>70</sup> In summary, the author proposes a highly active OER catalyst driven by a LOM with longterm stability.

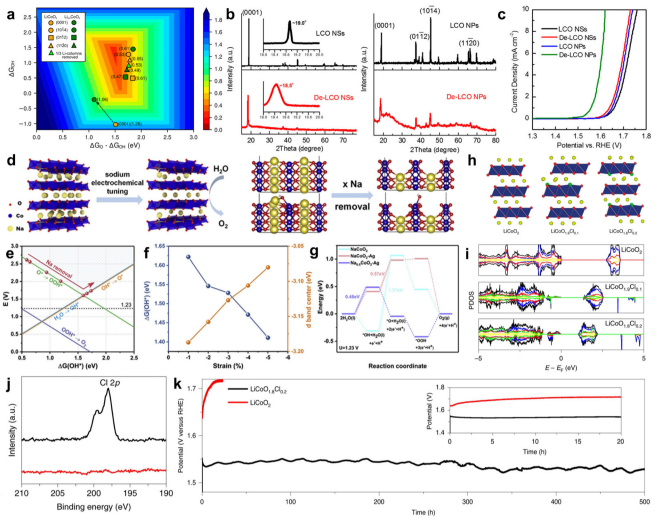
Cao *et al.* used metal-organic frameworks (MOFs) as precursors and utilized their multi-layer structure to peel MOFs into a single-layer structure through  $Ir^{3+}$ -assisted stripping, while doping Ir into its lattice. Then, Ir-Co<sub>3</sub>O<sub>4</sub> nanosheets (Ir-Co<sub>3</sub>O<sub>4</sub> NSs) were obtained by oxidation in air (Fig. 3k). It reduces the loading capacity of Ir, and it also exploits the interaction between metal and carrier to construct special structures. Density functional theory calculations indicate that the intermediate \*OOH in the OER process can stabilize by combining with lattice oxygen around the Ir active site through the formation of hydrogen bonds, greatly reducing the \*OOH formation energy of RDS and reducing the energy consumption of the OER process. The catalyst exhibits excellent water oxidation activity and stability. At 10 mA cm<sup>-2</sup>, the overpotential of this catalyst drops to 226 mV, which is much better than that of commercial IrO<sub>2</sub>. Meanwhile, the catalyst exhibits excellent corrosion resistance under acidic OER conditions, with a service life of up to 500 hours at 10 mA cm<sup>-2</sup>.<sup>77</sup> Layered single metal oxides offer the advantage of simplicity and ease of acquisition, rendering them exemplary materials for investigating the relationship between catalyst performance and mechanism. Nevertheless, striking a balance between the utilization of precious metals and catalytic efficacy often poses a formidable challenge. Hence, the imperative to develop other structured LMOs emerges.

#### 4.2 Layered alkali cobalt oxides

Alkali cobalt oxides such as LiCoO2 and NaCoO2, as types of alkali metal oxides, are natural layered structure materials composed of CoO<sub>2</sub> layers and alternating arrangement of Li<sup>+</sup> or Na<sup>+</sup> ions embedded in the layers. Therefore, they have flexible ion removal and insertion regulation and are commonly used in alkaline lithium-ion batteries. In recent years, the universality of electrochemical reactions has made it a new type of electrocatalyst in oxygen-related reactions.<sup>78,79</sup> However, compared to noble metal catalysts, their intrinsic activity is far from sufficient, so researchers have modified them through certain means to achieve high activity. Usually, by adjusting the insertion and extraction of Li<sup>+</sup> or Na<sup>+</sup> ions, the aim of adjusting the materials' bulk composition, crystal surface exposure, stress/strain, and coordination environment is tried to be achieved which is expected to improve the OER activity.<sup>80-82</sup> For example, due to the influence of exposed surface types on OER activity, Cui *et al.* obtained  $\text{Li}_x\text{CoO}_2$  ( $x \approx$ 0.5, De-LCO) by delithiation of LiCoO<sub>2</sub> (LCO) by an electrochemical method. Due to the high crystallinity of LCO obtained through the delithiation process with multiple surface exposures, it is interesting to study the effects of different surfaces and sites of LCO and De-LCO on OER activity. The author indicates through theoretical calculations that  $(11\overline{2}0)$  and  $(01\overline{1}2)$  are active surfaces, while (0001) is an inert surface (Fig. 4a). This may be attributed to an increase in the concentration of Co<sup>4+</sup> sites and a shift in the 2p state of reactive oxygen species. In order to confirm from experimental data, the author synthesized two different forms of LCO through experiments. From the XRD spectrum (Fig. 4b), it can be seen that LCO nanosheets (LCO NSs) are mainly exposed on the substrate (0001), while the substrate of LCO nanoparticles (LCO NPs) is significantly reduced to expose more edge sites. Subsequently, electrochemical delithiation was performed on both samples and the OER activity of crystalline LCO NSs was negligible, while the OER activity of LCO NPs was significantly enhanced (Fig. 4c). Combined with theoretical predictions, these experimental results confirm that the base surface of De-LCO is very inactive for OERs, while other major surfaces, in particular (11 $\overline{2}0$ ) and (01 $\overline{1}2$ ), are active, which is conducive to the OER process.80

In addition, Dai *et al.* used an electrochemical desorption process to remove sodium from layered NaCoO<sub>2</sub>, resulting in

**Dalton Transactions** 



**Fig. 4** (a) Theoretical calculation of the OER activity of the  $LiCOO_2/Li_{0.5}COO_2$  system (theoretical overpotential value). The arrow indicates the change in activity after delithiation. (b) XRD spectra of LCO NSs and LCO NPs before and after delithiation. The illustration shows a local fine XRD pattern. (c) Polarization curves of LCO NSs and LCO NPs before and after delithiation. Adapted with permission from ref. 80. Copyright 2017, American Chemical Society. (d) Schematic diagram of the electrochemical removal of Na<sup>+</sup>. NaCoO<sub>2</sub>(100) surface atomic model. (e) Free energy of \*OH corresponds to different degrees of Na<sup>+</sup> release from NaCoO<sub>2</sub>. (f) Relationship of the \*OH and d-band center free energy of NaCoO<sub>2</sub> vs. compressive strain. (g) Free energy diagram of the OER process at U = 1.23 V. Adapted with permission from ref. 66. Copyright 2021, Elsevier. (h) Schematic diagram of the atomic structure; Li, yellow; Co, blue; O, Red; Cl, green. (i) PDOS with total (black) and Li, yellow; Co, blue; O, Red; Cl, green. The PDOS of Li and Cl has been magnified by 20 times. (j) Cl 2p XPS: LiCoO<sub>1.8</sub>Cl<sub>0.2</sub> (black) and LiCoO<sub>2</sub> (red). (k) Stability at a current density of 20 mA cm<sup>-2</sup>. Adapted with permission from ref. 83. Copyright 2021, Springer Nature.

sodium-deficient Na<sub>0.7</sub>CoO<sub>2</sub> with a certain strain (Fig. 4d). Using  $\Delta G_{*OH}$  as the catalytic activity indicator for the OER process, the author calculated the Gibbs free energy values of the \*OH step under different degrees of desodiation. When the degree of desodiation is at a specific value, there is an optimal Gibbs free energy value (Fig. 4e). The desorption process of Na<sup>+</sup> corresponds to a decrease in lattice spacing and the occurrence of strain. Analysis shows that when the strain level is less than 5%, as the strain increases,  $\Delta G_{*OH}$  decreases and the Co d-band center shifts upwards, indicating an increase in the OER activity (Fig. 4f). Desodiation process has a few improvement in the OER process. Therefore, synchronous Ag<sup>+</sup> doping was carried out on this basis. Compressive strain was introduced within the layer while introducing tensile strain between the layers. Through DFT calculations, it was shown that the RDS Gibbs free energy of the catalyst doped with Ag<sup>+</sup> and desodiation significantly decreased, greatly improving the OER activity (Fig. 4g). Among them, the overpotential of Ag-Na<sub>0.7</sub>CoO<sub>2</sub> is the lowest 236 mV at 10 mA cm<sup>-2</sup>, with a minimum Tafel slope of 48 mV dec<sup>-1.66</sup> This work provides an interesting approach for interlayer/intralayer modification of layered materials to achieve efficient OERs. Lim *et al.* prepared a layered LiCoO<sub>2-x</sub>Cl<sub>x</sub> (x = 0, 0.1 or 0.2) by a solid-phase reaction method, and its atomic structure diagram is shown in Fig. 4h. DFT calculations indicate that Cl doping causes the d-band center of the active site to shift towards the

Fermi level and the gap between the valence and conduction bands decreases from 2.7 eV of LiCoO<sub>2</sub> to 1.3 eV and 1.0 eV of LiCoO<sub>1.9</sub>Cl<sub>0.1</sub> and LiCoO<sub>1.8</sub>Cl<sub>0.2</sub>, respectively (Fig. 4i). This indicates that Cl doping enhances the bulk conductivity, which is attributed to a decrease in electron transfer resistance. The author further confirmed the successful doping of Cl by comparing LiCoO<sub>2</sub> before and after Cl doping by XPS (Fig. 4j). The doping of Cl reduced the potential for in situ cobalt oxidation and lithium leaching, leading to the transformation of the LiCoO<sub>1.8</sub>Cl<sub>0.2</sub> surface into an amorphous (oxygen) hydroxide phase (CoOOH) during the OER process. In contrast, Cl-free LiCoO<sub>2</sub> requires a higher electrochemical potential to initiate surface reconstruction and requires long-term cycling to stabilize it. Therefore, the performance of surface recombination LiCoO<sub>1.8</sub>Cl<sub>0.2</sub> is superior to many state-of-the-art OER catalysts and exhibits significant stability, which can be maintained for 500 hours at a current density of 20 mA cm<sup>-2</sup> (Fig. 4k).<sup>83</sup> This work provides guidance for designing excellent OER catalysts by controlling in situ leaching of layered materials to regulate surface recombination.

#### 4.3 Perovskites

Perovskites are a large class of oxides, commonly known as ABO<sub>3</sub>, where A is usually a tombarthite or alkaline metal and B is a transition metal.<sup>84</sup> An attractive advantage of perovskite oxides is their high tunability.85,86 However, perovskite oxides typically exhibit cationic leaching behavior, which can lead to the collapse of the perovskite framework during the OER process, greatly reducing the stability of the bulk.<sup>87</sup> In addition to having a typical cubic structure, the perovskite family also has a layered structure. The flexibility of this layered perovskite in the configuration also contributes to its tunable electronic structure and stable IrO<sub>6</sub> framework.<sup>88</sup> This provides a prerequisite for improving the catalytic activity and structural stability of OER catalysts. Sr<sub>2</sub>IrO<sub>4</sub>, as a layered perovskite oxide, has a higher mass activity than that of IrO<sub>2</sub>. However, there are still some unstable factors such as leaching behavior.<sup>89,90</sup> Therefore, scientists modify the OER by preparing monolayers, ion exchange, constructing heterostructures, and other forms to achieve the goal of optimizing OER performance. Threedimensional perovskite SrIrO<sub>3</sub> is an OER catalyst with excellent performance, but its stability is poor due to severe Sr leaching during the OER process. Cho et al. constructed a Sr<sub>2</sub>IrO<sub>4</sub>/ SrIrO<sub>3</sub> heterostructure (SrIrO) by modifying 6H phase SrIrO<sub>3</sub> with layered Sr<sub>2</sub>IrO<sub>4</sub>. Layered Sr<sub>2</sub>IrO<sub>4</sub> is different from the traditional 3D structured perovskite. During the OER process, Sr<sup>+</sup> cations between layers can be easily leached while maintaining the IrO<sub>6</sub> octahedral framework with shared in-plane angles (Fig. 5a). In this way, both the bulk structure can be stabilized and oxygen can be transported through gaps. In order to better illustrate the structural evolution during the OER process, XPS analysis of SrIrO was performed in 0.1 M HClO<sub>4</sub> before and after electrocatalysis for 2 hours. The Sr 3d XPS spectrum is shown in Fig. 5b. Compared to the initial state, the Sr 3d peak of the catalyst significantly weakened after testing, indicating that a large amount of Sr<sup>+</sup> leaching formed

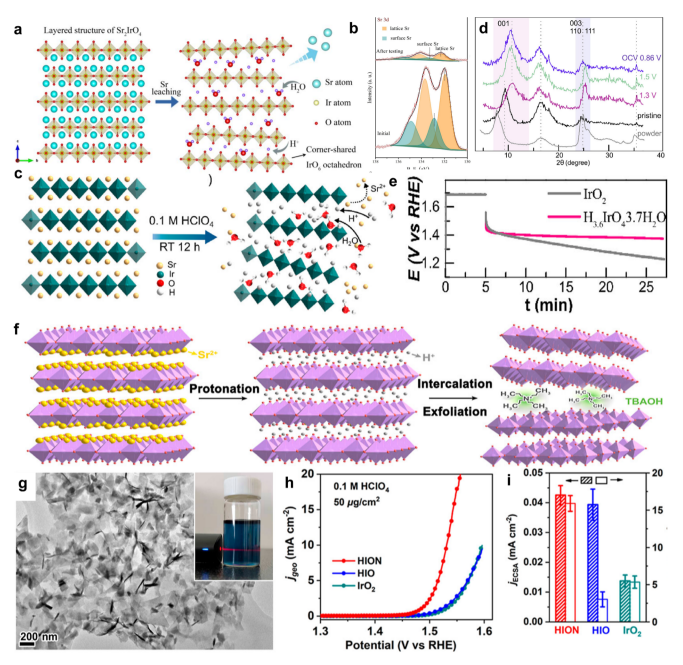
a Sr deficient surface after electrochemical testing. This further proves the leaching of Sr<sup>+</sup> cations through experiments. In addition, Sr leaching can lead to significant surface coarsening, thereby increasing the active surface area and affecting catalytic performance. SrIrO exhibits extremely low overpotential, with an OER of 245 mV at 10 mA cm<sup>-2</sup> and excellent catalytic stability in 0.5 M  $H_2SO_4$ .<sup>91</sup>

Grimaud *et al.* modified layered perovskite Sr<sub>2</sub>IrO<sub>4</sub> by a Sr<sup>+</sup>/ H<sup>+</sup> cation exchange for the first time and prepared the protonated phase H<sub>3.6</sub>IrO<sub>4</sub>·3.7H<sub>2</sub>O at room temperature (Fig. 5c). Usually, layered Sr<sub>2</sub>IrO<sub>4</sub> undergoes structural collapse of composite oxides during cationic leaching under acidic conditions, resulting in poor overall catalytic stability. Charge compensation via proton exchange helps avoid the formation of soluble Ir<sup>3+</sup> or IrO<sub>4</sub><sup>2-</sup>, improving stability. The author further demonstrated the occurrence of Sr<sup>+</sup>/H<sup>+</sup> cation exchange and the generation of protonated phase H<sub>3.6</sub>IrO<sub>4</sub>·3.7H<sub>2</sub>O through ex situ XRD at different OER process potentials. As shown in Fig. 5d, when scanning from an open circuit voltage to 1.3 V vs. RHE, the characteristic peak shifts towards a high angle. This indicates lattice shrinkage, and the volume shrinkage is much greater than the volume reduction caused by Ir<sup>4+</sup> oxidation, leading to Ir-O bond shortening, thus attributed to deprotonation and structural water in H<sub>3.6</sub>IrO<sub>4</sub>·3.7H<sub>2</sub>O. In summary, this cation exchange method stabilizes the structure of the layered perovskite material itself, giving it longer stability than that of commercial IrO2 (Fig. 5e).92 On the basis of cation exchange modification of layered perovskite Sr<sub>2</sub>IrO<sub>4</sub>, Zou et al. obtained protonated H<sub>4</sub>IrO<sub>4</sub> (HIO) by protonation treatment of layered perovskite Sr<sub>2</sub>IrO<sub>4</sub>, and, for the first time, obtained single- or few-layer protonated H<sub>4</sub>IrO<sub>4</sub> (HION) via liquid-phase exfoliation (Fig. 5f). The stripped HION is completely protonated, retaining the layered perovskite structure. The morphological integrity and high dispersibility of these protonated nanosheets result in extremely low Ir loading levels (30  $\mu$ g cm<sup>-2</sup>), and it exhibits excellent OER performance, with an activity 10 times that of IrO<sub>2</sub> (Fig. 5h). Through experimental results combined with theoretical analysis, it is shown that protonated perovskite nanosheets not only reduce Sr dissolution and structural reconstruction, but also expose more active sites. Moreover, layered perovskite frameworks have suitable electronic structures and efficient active sites to provide high intrinsic activity for OER processes (Fig. 5g). The activity of the catalyst was measured by an electrochemical active surface area (ECSA) method, and the results indicated that after normalization, HION and HIO had basically the same activity, much greater than IrO2, indicating a significant increase in the intrinsic activity of protonated layered perovskite (Fig. 5i).93

#### 4.4 Other natural layered metal oxides

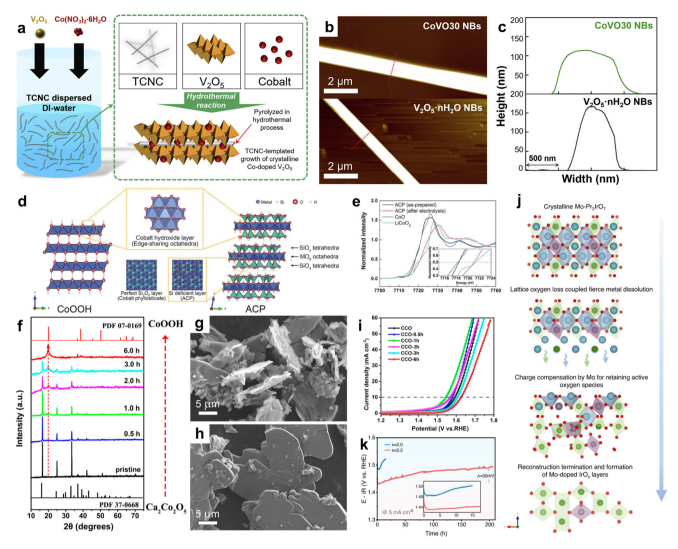
**4.4.1**  $V_2O_5 \cdot nH_2O$ . Vanadium oxide is a rich compound on Earth that has been extensively studied due to its unique physical and chemical properties.<sup>94–96</sup> In particular, hydrated vanadium pentoxide ( $V_2O_5 \cdot nH_2O$ ) has a layered structure, which is composed of  $V_2O_5$  and  $H_2O$  arranged alternately.<sup>97</sup>

#### Frontier



**Fig. 5** (a) Schematic diagram of the layered structure of  $Sr_2IrO_4$  and the structure after Sr leaching. (b) XPS spectra of Sr 3d for SrIrO before and after 2 hours of electrocatalysis. Adapted with permission from ref. 91. Copyright 2021, Elsevier. (c) Schematic diagram of  $Sr_2IrO_4$  and  $H_{3.6}IrO_4$ ·3.7 $H_2O$  structures. (d) *Ex situ* XRD patterns of  $H_{3.6}IrO_4$ ·3.7 $H_2O$  collected at different OER potentials. (e) Stability curve at a constant potential of 1.65 V vs. RHE. Adapted with permission from ref. 92. Copyright 2020, American Chemical Society. (f) Schematic diagram of  $Sr_2IrO_4$  liquid-phase exfoliation. Sr, yellow; O, red; H, gray;  $IrO_6$ , purple. (g) TEM image of HION. The illustration shows the dispersion of HION in water, exhibiting The Tyndall effect. (h) Polarization curve of HION in 0.1 M HClO<sub>4</sub>. (i) ECSA and ECSA-normalized specific activity of HION at 1.53 V vs. RHE. Adapted with permission from ref. 93. Copyright 2022, American Chemical Society.

The high accessibility of metal ions between two  $V_2O_5$  crystal layers suppresses vanadium dissolution or irreversible phase transitions in  $V_2O_5 \cdot nH_2O$ , making it highly adjustable and possible to achieve ideal stability.<sup>98,99</sup> Yoon *et al.* prepared Co single atom–doped  $V_2O_5 \cdot nH_2O$  layered nanobelts (CoVO NBs) using an organic template-assisted crystal growth method (Fig. 6a). The dispersed Co atoms and highly crystalline  $V_2O_5{\cdot}nH_2O$  promote the exposure of active sites and efficient charge and mass transfer of intermediate substances, resulting in excellent OER activity. The DFT calculation indicates that Co atoms can be doped at lattice and interlayer positions, thereby reducing interlayer spacings, weakening intermediate binding energy and reducing the reaction energy barrier of OERs. The author also indirectly explained this phenomenon



**Fig. 6** (a) Schematic diagram of the synthesized CoVO NBs. (b) AFM image. (c) AFM measures the width and thickness of  $V_2O_5 \cdot nH_2O$  NBs and CoVO30 NBs. Adapted with permission from ref. 100. Copyright 2022, Elsevier. (d) Schematic diagram of CoOOH and ACP structures. (e) XANES Co *k*-edge spectra of ACP before and after CV cycling. For comparison, CoO and LiCoO<sub>2</sub> spectra display reference Co<sup>2+</sup> and Co<sup>3+</sup>, respectively. Adapted with permission from ref. 43. Copyright 2017, WILEY. (f) XRD spectra of the original CCO and CCO-*x*h treated at different times. (g and h) SEM images of CCO-1h and CCO. (i) Polarization curves of COO and COO-*x*h. Adapted with permission from ref. 106. Copyright 2022, American Chemical Society. (j) Schematic diagram of electrochemical surface reconstruction process of Mo-Pr<sub>3</sub>IrO<sub>7</sub>. (k) Chronopotentiometry curves at 5 mA cm<sup>-2</sup> in 0.1 M HClO<sub>4</sub> electrolyte. Adapted with permission from ref. 107. Copyright 2023, Springer Nature.

from an experimental perspective. As shown in Fig. 6b and c. Atomic force microscopy (AFM) testing was conducted on the CoVO30 NBs (30 represents a certain doping amount of Co) and  $V_2O_5 \cdot nH_2O$  NBs after Co doping of initial samples. The results indicated that the thickness of the doped samples decreased significantly, indicating a decrease in interlayer spacing. The results showed that CoVO NBs exhibit a higher OER catalytic activity in 0.1 M KOH than that of the existing IrO<sub>2</sub> catalysts (Table 1). Despite the partial leaching of  $V^{4+}$ during the OER process, the Co group on the oxide skeleton still keeps high OER activity to maintain stability.<sup>100</sup>

**4.4.2 Cobalt phyllosilicate.** Research has shown that the coordination number of transition metals  $(MO_x)$ , their local distortions, and their connectivity properties can sensitively

affect catalytic activity.<sup>105–107</sup> In addition, adjacent redox inert elements can alter the redox activity of the catalyst. For example, anions such as phosphates and borates promote proton coupling and electron transfer.<sup>108,109</sup> Inspired by this, Kang *et al.* prepared a layered crystalline motif amorphous cobalt phyllosilicate (ACP) by a room-temperature precipitation method. The model crystal structure of ACP has the molecular formula  $Co_3Si_3O_6(OH)_6$ , characterized by: (i) a basic cobaltbased silicate structure, (ii) the presence of silanol groups originating from Si vacancies, and (iii) a Co/Si/O atomic ratio of 1/ 1/4. Compared to CoOOH, except for the silicate groups between  $CoO_6$  layers that only exist in ACP, all other layers are a series of  $CoO_6$  layers (Fig. 6d). In order to demonstrate the active substances in the OER process, the author compared the

Table 1	Electrocatalytic OEF	R performance of la	ayered metal oxide o	atalysts
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	Catalyst	Electrolyte	Overpotential at specific current density	Tafel slope (mV dec <sup>-1</sup> )	Stability at specific current density	Ref.
Layered single metal oxides	RuO <sub>2</sub> NSs	$0.5 \text{ M H}_2\text{SO}_4$	199 mV at 10 mA $cm^{-2}$	38.2	5.5 h at 10 mA $cm^{-2}$	73
, ,	1T-IrO <sub>2</sub>	$0.1 \text{ M HClO}_{4}$	197 mV at 10 mA $cm^{-2}$	49	126 h at 250 mA $cm^{-2}$	74
	Ir-MnO <sub>2</sub>	$0.5 \text{ M H}_2 \text{SO}_4$	218 mV at 10 mA cm <sup>-</sup>	59.61	650  h at 10 mA cm <sup>-2</sup>	70
	Ir-Co <sub>3</sub> O <sub>4</sub> NSs	$0.5 \text{ M H}_2^2 \text{SO}_4$	226 mV at 10 mA cm <sup>-</sup>	74.4	500 h at 10 mA $cm^{-2}$	77
MCoOx	De-Li <sub>r</sub> CoO <sub>2</sub>	0.1 M KOH	$390 \text{ mV}$ at $10 \text{ mA} \text{ cm}^{-2}$	57	$10 \text{ h}$ at $10 \text{ mA cm}^{-2}$	80
	Ag-Na <sub>0 7</sub> CoO <sub>2</sub>	1 M KOH	$236 \text{ mV}$ at $10 \text{ mA} \text{ cm}^{-2}$	59.2	$30 \text{ h}$ at $10 \text{ mA cm}^{-2}$	66
	LiCoO <sub>1 8</sub> Cl <sub>0 2</sub>	1 M KOH	$290 \text{ mV}$ at $40 \text{ mA} \text{ cm}^{-2}$	55.4	500 h at 20 mA cm <sup>-2</sup>	83
Sr <sub>2</sub> IrO <sub>4</sub>	Sr <sub>2</sub> IrO <sub>4</sub> /SrIrO <sub>3</sub>	0.5 M H <sub>2</sub> SO <sub>4</sub>	245  mV at 10 mA cm <sup>-2</sup>	47.4	40 h at 10 mA $cm^{-2}$	91
2 .	$H_{36}IrO_{4}\cdot 3.7H_{2}O$	$0.5 \text{ M H}_2 \text{SO}_4$	$300 \text{ mV}$ at $100 \text{ A g}_{\text{Ir}}^{-1}$	_	0.5 h at 1.65 V vs. RHE	92
	Single layered H <sub>4</sub> IrO <sub>4</sub>	0.1 M HClO <sub>4</sub>	300  mV at 10 mA cm <sup>-2</sup>	46.9	$30 \text{ h}$ at $10 \text{ mA cm}^{-2}$	93
	Co SACs- $V_2O_5 \cdot nH_2O$	0.1 M KOH	428  mV at 20 mA cm <sup>-2</sup>	92	10  h at 20 mA cm <sup>-2</sup>	100
	Co <sub>3</sub> Si <sub>3</sub> O <sub>6</sub> (OH) <sub>6</sub>	1 M KOH	$367 \text{ mV}$ at $10 \text{ mA} \text{ cm}^{-2}$	60	24  h at 10 mA cm <sup>-2</sup>	43
	Ca <sub>2</sub> Co <sub>2</sub> O <sub>5</sub> /CoOOH	1 M KOH	$320 \text{ mV}$ at $10 \text{ mA} \text{ cm}^{-2}$	91	$30 \text{ h}$ at $10 \text{ mA cm}^{-2}$	101
	$Pr_3Ir_{1-r}Mo_rO_7$	0.1 M HClO <sub>4</sub>	259  mV at 10 mA cm <sup>-2</sup>	50.42	200  h at 5 mA cm <sup>-2</sup>	102
	Commercial IrO <sub>2</sub>	$0.5 \text{ M H}_2 \text{SO}_4$	$397 \text{ mV}$ at $10 \text{ mA} \text{ cm}^{-2}$	142.4	4 h at 20 mA cm <sup><math>-2</math></sup>	103
	2	1 M KOH	386 mV at 10 mA $\mathrm{cm}^{-2}$	152.77	about 4 h at 10 mA $\rm cm^{-2}$	104

samples before and after the ACP electrocatalytic reaction using X-ray absorption near edge structure (XANES) (Fig. 6e). The valence of Co transferred from  $2^+$  to  $3^+$  before and after the test, indicating an irreversible change in Co during the OER process, which is consistent with the CoOOH catalytic process. Therefore, the improvement in the performance of ACP compared to CoOOH can be attributed to the adjustment of the local environment of Co active sites by silicate groups, thereby reducing the overpotential of the reaction process and achieving excellent OER performance.<sup>43</sup> This discovery suggests that introducing redox inert groups into the layer space of metal (oxygen) hydroxides to regulate the structure may be a feasible strategy for adjusting the catalytic activity.

4.4.3 Brownmillerite. The true active site of Co-based catalysts is usually attributed to cobalt oxyhydroxide (CoOOH), which is usually generated in situ on the surface of the electrocatalyst under OER conditions.<sup>110–112</sup> However, directly preparing pure phase CoOOH has poor activity during the OER process.<sup>113-115</sup> Therefore, how to prepare pre-catalysts reasonably, increase the exposure of active sites, and improve the intrinsic activity of active sites has become the fundamental starting point for the design of Co-based catalysts. Sun et al. prepared CoOOH (CCO-xh, x means time) with different thicknesses grown on the surface of brownmillerite Ca<sub>2</sub>Co<sub>2</sub>O<sub>5</sub> (CCO) using surface acid etching method (degree varies with etching time). Brownmillerite has a layered structure, which has the advantage of accurately controlling the surface structure, providing a good foundation for understanding the structural activity relationship between surface cobalt oxide and the substrate. From the XRD spectrum (Fig. 6f), it can be seen that with the extension of etching time, the brownmillerite undergoes a transition from the initial CCO phase to the final CoOOH phase. Through SEM observation (Fig. 6g and h), compared to CCO before etching, the sample after 1 hour of acid etching showed a fluffy state, with partial separation of layered CCO and a few layers of CoOOH appearing on the substrate surface. The experimental results indicate that the interface interaction of the composite catalyst is beneficial for promoting the insertion of hydrogen oxides, thereby increasing the involvement of lattice oxygen in OERs. By adjusting the surface structure reasonably, CCO-1h exhibits excellent OER activity. CCO-1h exhibits the best performance after 1 hour of etching, with an overpotential of 320 mV at 10 mA cm<sup>-2</sup> in 1 M KOH (Fig. 6i).<sup>101</sup> This work establishes a foundation for quantitatively explaining the relationship between the catalytic activity and the structure of active substances and substrates by utilizing the adjustability of layered materials.

4.4.4 Weberite. Weberite-type  $Ln_3IrO_7$  is an  $IrO_6$  octahedron with angular connections along the c-axis and a layered structure along the a-axis. Due to the O 2p orbital being near the Fermi level, the electrochemical process easily activates lattice oxygen redox.116 However, excessive involvement of lattice oxygen can easily lead to structural collapse and variable valence metals compensate for the intense leaching of Ln cations, effectively avoiding excessive loss of lattice oxygen and further dissolution of Ir.<sup>117</sup> For example, Zou et al. used high-valence Mo-modulated orthogonal Pr<sub>3</sub>Ir<sub>1-x</sub>Mo<sub>x</sub>O<sub>7</sub> as a model to activate lattice oxygen and cations, achieving directional and accelerated surface reconstruction. Doped Mo is not only used to optimize Ir-O covalence and make Pr easier to dissolve, thereby accelerating surface reconstruction, but also improves durability due to Mo buffering charge compensation, thereby preventing severe Ir dissolution and excessive lattice oxygen loss, ultimately maintaining the Mo-doped  $IrO_x$  layer as an active material. The structural diagram of its reconstruction process is shown in Fig. 6j. Therefore, Ir-O-Mo can be generated directionally, promoting the deprotonation of oxygen intermediates through the bridging oxygen-assisted deprotonation pathway. Therefore, the optimal catalyst exhibits the best activity with an overpotential of 259 mV, reaching 10 mA cm<sup>-2</sup>, 50 mV lower than the undoped counterpart, and exhibits better stability over 200 hours (Fig. 6k).<sup>102</sup> This work provides a directed surface reconstruction strategy for constructing strong Brønsted acid sites in  $IrO_x$  species, showcasing the potential for targeted electrocatalyst manufacturing under in situ actual reaction conditions.

## 5. Conclusion and perspectives

In summary, to secure the pivotal role of electrolytic water hydrogen production in the forthcoming energy landscape, the engagement of the OER process is imperative. Because the OER process involves a four-electron transfer step, which requires overcoming large reaction potential barriers and slow kinetics. Investigating novel, efficient, and cost-effective OER electrocatalysts to attain reduced energy consumption in the overall electrolysis of water has emerged as a pivotal research endeavor. LMOs inherently possess natural structural and compositional advantages. Layered architecture has the capability to augment both the specific surface area and the array of active behaviors. Compositionally, metal oxides can function as precursors, undergoing transformation into genuinely active species such as "OOH". Consequently, investigating layered metal oxide materials as the focal point of OER catalyst research facilitates elucidating the interplay among catalytic properties, mechanisms, and material architectures, thereby enabling the scientific design of electrolytic water catalysts. This article presents the inaugural overview of the recent application of catalysts based on LMOs in the field of OER catalysts. Layered metal oxide-based catalysts are categorized into four groups: single metal oxide (monolayer or few-layer structures acquired through specialized methodologies), alkali cobalt oxide, perovskite, and other metal oxides. The article consolidates the insights into structural advantages, design methodologies, catalytic mechanisms, performance characteristics, and other pertinent facets. The focus is on the necessary connection between the inherent mechanism of catalytic processes and the effective improvement of catalytic properties. For the intrinsic activity of layered metal oxide catalysts, within the framework of the AEM, one crucial consideration entails the adjustment of the d-band center of the transition metal active site to promote the adsorption and desorption of intermediates. In addition, adjustments to the electronic state surrounding oxygen atoms within the catalyst matrix are required to initiate both LOM and OPM while addressing the potential for structural collapse. Ultimately, the overarching objective is to strike a delicate balance between catalytic activity and structural stability.

To sum up, the catalytic activity is intricately related to the morphology, composition, coordination environment of active sites, electronic structure, *etc.*, of materials, whereas catalytic stability is closely associated with the coordination environment, chemical bonds, *etc.*, of materials. Despite significant endeavors by researchers in this domain, there remain unresolved issues necessitating thorough investigation in the field of universal design and mechanistic research of OER catalysts.

#### 5.1 Challenges

Although many studies have been conducted on LMOs in the field of OER, there are still many problems that need to be solved: (i) for non-natural layered materials, the method of preparing single-layer or few-layer catalysts is relatively limited, which cannot achieve widespread application. (ii) Conversely, the structural singularity of natural LMOs offers little reference utility for other layered materials, fostering inadvertent design outcomes, resulting in accidental design. (iii) A lack of standardized methodologies for assessing the stability of LMOs exacerbates the challenges within the field, and only a few studies exploring the LOM are involved. (iv) Typically confined to laboratory settings, the application of LMOs in OERs has not yet achieved commercial high-current applications.

#### 5.2 Development direction

5.2.1. Descriptor for catalyst design. Despite numerous studies employing diverse methodologies on LMOs, significant gaps persist in our understanding of the fundamental chemical properties of catalysts. The development of a singular or standardized descriptor grounded in the fundamental physical and chemical properties of catalyst materials holds the potential to circumvent traditional trial-and-error approaches, enabling a systematic catalyst design through scientific methodologies. For example, our analysis revealed that in traditional AEM, the adsorption energy of intermediates typically serves as a descriptor for predicting catalytic performance, often correlated with the electronic structure.<sup>81,118,119</sup> However, during the actual experimental procedure, it is also influenced by diverse factors including lattice spacing, bonding length, and the exposure of distinct crystal planes of the catalytic materials.<sup>66,70,80</sup> Therefore, it is necessary to search for a suitable unified descriptor in LMOs to study universal OER catalysts.

5.2.2. Relationship between catalytic mechanism and catalytic activity. Mastering the relationship between catalytic activity and catalytic mechanism in the OER process is pivotal for the design of basic materials for efficient OER catalysts. The characterization of XPS, XANES and theoretical calculations has been extensively employed for the research of the oxidation state, geometry, and electronic structure of various catalysts in the OER process.<sup>120-122</sup> However, the primary factors conducive to enhancing the catalytic process have yet to be precisely identified. Particularly for layered metal oxidebased materials, their high specific surface area is relatively large, and changes in the coordination environment are introduced in subsequent modifications. This requires the utilization of various characterization methods and post-processing methods to accurately conclude the main factors contributing to increase activity when investigating the catalytic mechanism for performance improvement. This serves as a scientific foundation for the rational design of catalysts.

**5.2.3.** Commercial application. The current application of LMOs in OER catalysis is confined to the laboratory testing phase at low current, such as 10 mV cm<sup>-2</sup> and 20 mV cm<sup>-2</sup>.<sup>2,123,124</sup> There are scarce instances of their application in the assembly of electrolytic cells or larger-scale equipment. Despite some studies confirming, *via* experimental and theoretical approaches, that specific coordination environments or charge compensation mechanisms can uphold long-term stability, their implementation at industrial current densities remains elusive. In addition, the lack of standardized activity

#### Frontier

evaluation and comparison schemes makes strict evaluation of electrocatalysts extremely difficult. Thus, it is imperative to establish unified standards and corresponding parameters for industrial application with regard to catalyst activity and stability testing, aiming to accelerate the commercialization process.

5.2.4 Drawing inferences from one example. At present, varieties of modification techniques exist for LMOs, encompassing defect engineering, phase engineering, doping, heterojunction formation, and other primary methods. In fact, a rational modification approach can theoretically be extrapolated for the modification of other LMOs. For example, Kang et al. achieved the design of an inert coordination environment for Co active sites by embedding defective SiO<sub>4</sub> ligands between layered CoO<sub>6</sub> structures, thereby improving the surrounding electronic structure of Co sites, which tends to favor the OER process.<sup>43</sup> Similarly, this method can be extended to the interlayers of alternative layered materials, enabling the prediction and realization of enhanced OER performance via theoretical and experimental approaches. Therefore, the design of catalysts in the future should embrace universality, integrating diverse methods with a range of structural materials and forecasting the performance of a given catalyst on a large scale to realize the objective of efficient catalyst screening.

# Conflicts of interest

There are no conflicts to declare.

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# References

- 1 H. Yang, F. Li, S. Zhan, Y. Liu, W. Li, Q. Meng, A. Kravchenko, T. Liu, Y. Yang and Y. Fang, *Nat. Catal.*, 2022, 5, 414–429.
- 2 X.-L. Zhang, P.-C. Yu, X.-Z. Su, S.-J. Hu, L. Shi, Y.-H. Wang, P.-P. Yang, F.-Y. Gao, Z.-Z. Wu and L.-P. Chi, *Sci. Adv.*, 2023, **9**, eadh2885.
- 3 L. E. Heim, D. Thiel, C. Gedig, J. Deska and M. H. Prechtl, *Angew. Chem., Int. Ed.*, 2015, 54, 10308–10312.
- 4 Y. Liu, L. Li, L. Wang, N. Li, X. Zhao, Y. Chen, T. Sakthivel and Z. Dai, *Nat. Commun.*, 2024, **15**, 2851.
- 5 N. Kittner, F. Lill and D. M. Kammen, *Nat. Energy*, 2017, 2, 1–6.
- 6 J. Zhang, Q. Zhang and X. Feng, *Adv. Mater.*, 2019, **31**, 1808167.

- 7 F. Jiao, J. Wang, Y. Lin, J. Li, X. Jing and Y. Gong, *Appl. Surf. Sci.*, 2021, 553, 149440.
- 8 H. Shi, X.-Y. Sun, S.-P. Zeng, Y. Liu, G.-F. Han, T.-H. Wang,
   Z. Wen, Q.-R. Fang, X.-Y. Lang and Q. Jiang, *Small Struct.*,
   2023, 4, 2300042.
- 9 Y. A. Situmorang, Z. Zhao, P. An, T. Yu, J. Rizkiana, A. Abudula and G. Guan, *Appl. Energy*, 2020, **268**, 115122.
- 10 U. Izquierdo, S. Neuberg, S. Pecov, H. Pennemann, R. Zapf, M. Wichert, V. Barrio, J. Cambra and G. Kolb, *Chem. Eng. J.*, 2017, **313**, 1494–1508.
- 11 L. Ouyang, J. Jiang, K. Chen, M. Zhu and Z. Liu, *Nano-Micro Lett.*, 2021, **13**, 134.
- 12 T. Wang, L. Tao, X. Zhu, C. Chen, W. Chen, S. Du, Y. Zhou, B. Zhou, D. Wang and C. Xie, *Nat. Catal.*, 2022, 5, 66–73.
- 13 X. Yan, M. Xia, H. Liu, B. Zhang, C. Chang, L. Wang and G. Yang, *Nat. Commun.*, 2023, 14, 1741.
- 14 Y. Chen, Y. Liu, L. Li, T. Sakthivel, Z. Guo and Z. Dai, *Adv. Funct. Mater.*, 2024, 2401452.
- 15 Y. Jiao, Y. Zheng, M. Jaroniec and S. Z. Qiao, *Chem. Soc. Rev.*, 2015, 44, 2060–2086.
- 16 W. Zhai, Y. Chen, Y. Liu, Y. Ma, P. Vijayakumar, Y. Qin, Y. Qu and Z. Dai, *Nano-Micro Lett.*, 2024, **16**, 115.
- 17 L. Yao, D. Wei, D. Yan and C. Hu, *Chem. Asian J.*, 2015, **10**, 630–636.
- 18 Y. Chen, Y. Liu, W. Zhai, H. Liu, T. Sakthivel, S. Guo and Z. Dai, *Adv. Energy Mater.*, 2024, 2400059.
- 19 A. Ali, F. Long and P. K. Shen, *Electrochem. Energy Rev.*, 2022, 5, 1.
- 20 J. S. Kim, B. Kim, H. Kim and K. Kang, *Adv. Energy Mater.*, 2018, **8**, 1702774.
- 21 X. Xu, Y. Pan, L. Ge, Y. Chen, X. Mao, D. Guan, M. Li, Y. Zhong, Z. Hu and V. K. Peterson, *Small*, 2021, 17, 2101573.
- 22 Z. P. Wu, X. F. Lu, S. Q. Zang and X. W. Lou, Adv. Funct. Mater., 2020, 30, 1910274.
- 23 P. R. Chowdhury, H. Medhi, K. G. Bhattacharyya and C. M. Hussain, *Coord. Chem. Rev.*, 2023, **483**, 215083.
- 24 X.-W. Lv, W.-W. Tian and Z.-Y. Yuan, *Electrochem. Energy Rev.*, 2023, **6**, 23.
- 25 X. Huang, X. Xu, C. Li, D. Wu, D. Cheng and D. Cao, *Adv. Energy Mater.*, 2019, **9**, 1803970.
- 26 J. Wang, Y. Gao, H. Kong, J. Kim, S. Choi, F. Ciucci, Y. Hao, S. Yang, Z. Shao and J. Lim, *Chem. Soc. Rev.*, 2020, 49, 9154–9196.
- 27 Q. Wang, Z. Zhang, C. Cai, M. Wang, Z. L. Zhao, M. Li, X. Huang, S. Han, H. Zhou and Z. Feng, *J. Am. Chem. Soc.*, 2021, 143, 13605–13615.
- M. Li, X. Wang, K. Liu, H. Sun, D. Sun, K. Huang, Y. Tang,
   W. Xing, H. Li and G. Fu, *Adv. Mater.*, 2023, 35, 2302462.
- 29 S. Roy, P. Dahiya, T. K. Mandal and S. Roy, *Dalton Trans.*, 2024, **53**, 5484–5494.
- 30 H. Yan, Z. Jiang, B. Deng, Y. Wang and Z. J. Jiang, *Adv. Energy Mater.*, 2023, **13**, 2300152.
- 31 R. Lin, T. Lin, J. Huang, X. Huang and Y. Liu, *Electrochim. Acta*, 2018, **281**, 348–356.

- 32 J. Ban, H. Xu, G. Cao, Y. Fan, W. K. Pang, G. Shao and J. Hu, *Adv. Funct. Mater.*, 2023, 33, 2300623.
- 33 Y. Wang, Y. Jiao, H. Yan, G. Yang, C. Tian, A. Wu, Y. Liu and H. Fu, *Angew. Chem.*, 2022, **134**, e202116233.
- 34 Z. Jiang, W. Zhou, C. Hu, X. Luo, W. Zeng, X. Gong,
  Y. Yang, T. Yu, W. Lei and C. Yuan, *Adv. Mater.*, 2023, 35, 2300505.
- 35 Y. Cao, Y. Wen, Y. Li, M. Cao, B. Li, Q. Shen and W. Gu, *Dalton Trans.*, 2024, **53**, 5291–5300.
- 36 J.-Y. Luo, Y. Yuan, H.-Y. Ruan, X.-Q. Wu, Y.-P. Wu, S. Li, G. Zhang, S. Sun and D.-S. Li, *Small Struct.*, 2023, 4, 2300074.
- 37 M. Bachhav, G. Pawar, F. Vurpillot, R. L. Danoix,
  F. D. R. Danoix, B. Hannoyer, Y. Dong and E. Marquis,
  J. Phys. Chem. C, 2018, 123, 1313–1319.
- 38 T. C. Ozawa and S. M. Kauzlarich, *Sci. Technol. Adv. Mater.*, 2008, **9**, 033003.
- 39 D. Weber, L. M. Schoop, D. Wurmbrand, J. R. Nuss, E. M. Seibel, F. F. Tafti, H. Ji, R. J. Cava, R. E. Dinnebier and B. V. Lotsch, *Chem. Mater.*, 2017, 29, 8338–8345.
- 40 S. Hao, M. Liu, J. Pan, X. Liu, X. Tan, N. Xu, Y. He, L. Lei and X. Zhang, *Nat. Commun.*, 2020, **11**, 5368.
- 41 F. Geng, R. Ma, Y. Ebina, Y. Yamauchi, N. Miyamoto and T. Sasaki, *J. Am. Chem. Soc.*, 2014, **136**, 5491–5500.
- 42 M. E. Strayer, T. P. Senftle, J. P. Winterstein, N. M. Vargas-Barbosa, R. Sharma, R. M. Rioux, M. J. Janik and T. E. Mallouk, *J. Am. Chem. Soc.*, 2015, 137, 16216– 16224.
- 43 J. S. Kim, I. Park, E. S. Jeong, K. Jin, W. M. Seong, G. Yoon, H. Kim, B. Kim, K. T. Nam and K. Kang, *Adv. Mater.*, 2017, **29**, 1606893.
- 44 Q. Wang, K. Dastafkan and C. Zhao, *Curr. Opin. Electrochem.*, 2018, **10**, 16–23.
- 45 P. Ganter, L. M. Schoop and B. V. Lotsch, *Adv. Mater.*, 2016, **29**, 1604884.
- 46 S. Jin, Journal, 2017, 2, 1937–1938.
- 47 X. Cai, R. Ma, T. C. Ozawa, N. Sakai, A. Funatsu and T. Sasaki, *Nanoscale*, 2014, **6**, 14419–14427.
- 48 A. S. Gupta, H. Akamatsu, F. G. Brown, M. A. T. Nguyen, M. E. Strayer, S. Lapidus, S. Yoshida, K. Fujita, K. Tanaka, I. Tanaka, T. E. Mallouk and V. Gopalan, *Chem. Mater.*, 2017, 29, 656–665.
- 49 H. Ishaq, I. Dincer and C. Crawford, Int. J. Hydrogen Energy, 2022, 47, 26238–26264.
- 50 N. M. Dowell, N. Sunny, N. Brandon, H. Herzog, A. Y. Ku, W. Maas, A. Ramirez, D. M. Reiner, G. N. Sant and N. Shah, *Joule*, 2021, 5, 2524–2529.
- 51 X. Ren, L. Dong, D. Xu and B. Hu, *Int. J. Hydrogen Energy*, 2020, **45**, 34326–34345.
- 52 S. Zhiznin, V. Timokhov and A. Gusev, *Int. J. Hydrogen* Energy, 2020, 45, 31353-31366.
- 53 A. Grimaud, O. Diaz-Morales, B. Han, W. T. Hong, Y.-L. Lee, L. Giordano, K. A. Stoerzinger, M. T. Koper and Y. Shao-Horn, *Nat. Chem.*, 2017, 9, 457–465.
- 54 W. Zhai, Y. Chen, Y. Liu, T. Sakthivel, Y. Ma, Y. Qin, Y. Qu and Z. Dai, *ACS Nano*, 2023, **17**, 17254–17264.

- 55 M. Okamura, M. Kondo, R. Kuga, Y. Kurashige, T. Yanai,
  S. Hayami, V. K. Praneeth, M. Yoshida, K. Yoneda and
  S. Kawata, *Nature*, 2016, 530, 465–468.
- 56 Y. Liu, T. Sakthivel, F. Hu, Y. Tian, D. Wu, E. H. Ang, H. Liu, S. Guo, S. Peng and Z. Dai, *Adv. Energy Mater.*, 2023, 13, 2203797.
- 57 J. Shan, Y. Zheng, B. Shi, K. Davey and S.-Z. Qiao, ACS Energy Lett., 2019, 4, 2719–2730.
- 58 R. Gao, J. Zhu and D. Yan, *Nanoscale*, 2021, **13**, 13593–13603.
- 59 L. Li, P. Wang, Q. Shao and X. Huang, *Chem. Soc. Rev.*, 2020, **49**, 3072–3106.
- 60 N.-T. Suen, S.-F. Hung, Q. Quan, N. Zhang, Y.-J. Xu and H. M. Chen, *Chem. Soc. Rev.*, 2017, 46, 337–365.
- 61 L. Gao, X. Cui, C. D. Sewell, J. Li and Z. Lin, *Chem. Soc. Rev.*, 2021, **50**, 8428–8469.
- 62 T. Qiu, B. Tu, D. Saldana-Greco and A. M. Rappe, ACS Catal., 2018, 8, 2218–2224.
- 63 L. Hu, L. Wu, M. Liao, X. Hu and X. Fang, Adv. Funct. Mater., 2012, 22, 998–1004.
- 64 J. Suntivich, K. J. May, H. A. Gasteiger, J. B. Goodenough and Y. Shao-Horn, *Science*, 2011, 334, 1383–1385.
- 65 A. Vojvodic and J. K. Nørskov, Science, 2011, 334, 1355-1356.
- 66 L. Sun, Z. Dai, L. Zhong, Y. Zhao, Y. Cheng, S. Chong, G. Chen, C. Yan, X. Zhang and H. Tan, *Appl. Catal.*, *B*, 2021, 297, 120477.
- 67 Y. Pan, X. Xu, Y. Zhong, L. Ge, Y. Chen, J.-P. M. Veder, D. Guan, R. O'Hayre, M. Li and G. Wang, *Nat. Commun.*, 2020, **11**, 2002.
- 68 W. T. Hong, M. Risch, K. A. Stoerzinger, A. Grimaud, J. Suntivich and Y. Shao-Horn, *Energy Environ. Sci.*, 2015, 8, 1404–1427.
- 69 P. R. Chowdhury, H. Medhi, K. G. Bhattacharyya and C. M. Hussain, *Coord. Chem. Rev.*, 2024, **501**, 215547.
- 70 Z. Shi, Y. Wang, J. Li, X. Wang, Y. Wang, Y. Li, W. Xu,
   Z. Jiang, C. Liu and W. Xing, *Joule*, 2021, 5, 2164–2176.
- 71 T. Reier, H. N. Nong, D. Teschner, R. Schlögl and P. Strasser, *Adv. Energy Mater.*, 2017, 7, 1601275.
- 72 S. Yagi, I. Yamada, H. Tsukasaki, A. Seno, M. Murakami, H. Fujii, H. Chen, N. Umezawa, H. Abe and N. Nishiyama, *Nat. Commun.*, 2015, 6, 8249.
- 73 Z. L. Zhao, Q. Wang, X. Huang, Q. Feng, S. Gu, Z. Zhang,
  H. Xu, L. Zeng, M. Gu and H. Li, *Energy Environ. Sci.*, 2020, 13, 5143–5151.
- 74 Q. Dang, H. Lin, Z. Fan, L. Ma, Q. Shao, Y. Ji, F. Zheng, S. Geng, S.-Z. Yang and N. Kong, *Nat. Commun.*, 2021, 12, 6007.
- 75 S. Wang, T. Shen, C. Yang, G. Luo and D. Wang, *ACS Catal.*, 2023, **13**, 8670–8691.
- 76 S. Ge, R. Xie, B. Huang, Z. Zhang, H. Liu, X. Kang, S. Hu, S. Li, Y. Luo and Q. Yu, *Energy Environ. Sci.*, 2023, 16, 3734–3742.
- 77 Y. Liu, Y. Chen, X. Mu, Z. Wu, X. Jin, J. Li, Y. Xu, L. Yang,
   X. Xi and H. Jang, ACS Catal., 2023, 13, 3757–3767.
- 78 M. S. Islam, M. Kim, X. Jin, S. M. Oh, N.-S. Lee, H. Kim and S.-J. Hwang, ACS Energy Lett., 2018, 3, 952–960.

Published on 30 2567. Downloaded on 25/7/2567 9:54:39.

- 79 Z. Dai, U. Mani, H. T. Tan and Q. Yan, *Small Methods*, 2017, **1**, 1700098.
- Z. Lu, G. Chen, Y. Li, H. Wang, J. Xie, L. Liao, C. Liu, Y. Liu,
   T. Wu and Y. Li, *J. Am. Chem. Soc.*, 2017, 139, 6270–6276.
- 81 Z. Lu, H. Wang, D. Kong, K. Yan, P.-C. Hsu, G. Zheng, H. Yao, Z. Liang, X. Sun and Y. Cui, *Nat. Commun.*, 2014, 5, 4345.
- 82 Z. Lu, K. Jiang, G. Chen, H. Wang and Y. Cui, *Adv. Mater.*, 2018, **30**, 1800978.
- 83 J. Wang, S.-J. Kim, J. Liu, Y. Gao, S. Choi, J. Han, H. Shin, S. Jo, J. Kim and F. Ciucci, *Nat. Catal.*, 2021, 4, 212–222.
- 84 Y. Chen, H. Li, J. Wang, Y. Du, S. Xi, Y. Sun, M. Sherburne, J. W. Ager III, A. C. Fisher and Z. J. Xu, *Nat. Commun.*, 2019, **10**, 572.
- 85 R. Tang, Y. Nie, J. K. Kawasaki, D.-Y. Kuo, G. Petretto, G. Hautier, G.-M. Rignanese, K. M. Shen, D. G. Schlom and J. Suntivich, *J. Mater. Chem. A*, 2016, 4, 6831–6836.
- 86 A. Grimaud, A. Demortière, M. Saubanère, W. Dachraoui, M. Duchamp, M.-L. Doublet and J.-M. Tarascon, *Nat. Energy*, 2016, 2, 1–10.
- 87 H. J. Song, H. Yoon, B. Ju and D. W. Kim, Adv. Energy Mater., 2021, 11, 2002428.
- 88 C. Sun, J. A. Alonso and J. Bian, *Adv. Energy Mater.*, 2021, 11, 2000459.
- 89 Y. K. Kim, N. Sung, J. Denlinger and B. Kim, *Nat. Phys.*, 2016, 12, 37–41.
- 90 H. Tsai, W. Nie, J.-C. Blancon, C. C. Stoumpos, R. Asadpour, B. Harutyunyan, A. J. Neukirch, R. Verduzco, J. J. Crochet and S. Tretiak, *Nature*, 2016, **536**, 312–316.
- 91 L. Zhang, H. Jang, Z. Li, H. Liu, M. G. Kim, X. Liu and J. Cho, *Chem. Eng. J.*, 2021, **419**, 129604.
- 92 R. Zhang, P. E. Pearce, V. Pimenta, J. Cabana, H. Li, D. A. D. Corte, A. M. Abakumov, G. Rousse, D. Giaume and M. Deschamps, *Chem. Mater.*, 2020, **32**, 3499–3509.
- 93 H. Chen, L. Shi, K. Sun, K. Zhang, Q. Liu, J. Ge, X. Liang,
  B. Tian, Y. Huang and Z. Shi, ACS Catal., 2022, 12, 8658– 8666.
- 94 T. Wu, K. Zhu, C. Qin and K. Huang, J. Mater. Chem. A, 2019, 7, 5612–5620.
- 95 M. Liu, B. Su, Y. Tang, X. Jiang and A. Yu, Adv. Energy Mater., 2017, 7, 1700885.
- 96 A. S. Etman, H. D. Asfaw, N. Yuan, J. Li, Z. Zhou, F. Peng,
  I. Persson, X. Zou, T. Gustafsson and K. Edström, *J. Mater. Chem. A*, 2016, 4, 17988–18001.
- 97 D. Kundu, B. D. Adams, V. Duffort, S. H. Vajargah and L. F. Nazar, *Nat. Energy*, 2016, 1, 1–8.
- 98 K. Fan, H. Chen, Y. Ji, H. Huang, P. M. Claesson, Q. Daniel, B. Philippe, H. Rensmo, F. Li and Y. Luo, *Nat. Commun.*, 2016, 7, 11981.
- 99 A. Moretti, F. Maroni, I. Osada, F. Nobili and S. Passerini, *ChemElectroChem*, 2015, **2**, 529–537.
- 100 C. Youn, S. Shin, K. Shin, C. Kim, C.-L. Park, J. Choi, S. H. Kim, S. Y. Yeo, M. W. Shin and G. Henkelman, *Chem Catal.*, 2022, 2, 1191–1210.
- 101 S. Song, L. Mu, Y. Jiang, J. Sun, Y. Zhang, G. Shi and H. Sun, ACS Appl. Mater. Interfaces, 2022, 14, 47560–47567.

- 102 S. Chen, S. Zhang, L. Guo, L. Pan, C. Shi, X. Zhang, Z.-F. Huang, G. Yang and J.-J. Zou, *Nat. Commun.*, 2023, 14, 4127.
- 103 X. Zheng, J. Yang, P. Li, Q. Wang, J. Wu, E. Zhang, S. Chen, Z. Zhuang, W. Lai and S. Dou, *Sci. Adv.*, 2023, 9, eadi8025.
- 104 M. Hou, L. Zheng, D. Zhao, X. Tan, W. Feng, J. Fu, T. Wei, M. Cao, J. Zhang and C. Chen, *Nat. Commun.*, 2024, 15, 1342.
- 105 J. Park, H. Kim, K. Jin, B. J. Lee, Y.-S. Park, H. Kim, I. Park, K. D. Yang, H.-Y. Jeong and J. Kim, *J. Am. Chem. Soc.*, 2014, **136**, 4201–4211.
- 106 K. A. Stoerzinger, W. S. Choi, H. Jeen, H. N. Lee and Y. Shao-Horn, *J. Phys. Chem. Lett.*, 2015, **6**, 487–492.
- 107 A. Grimaud, C. E. Carlton, M. Risch, W. T. Hong, K. J. May and Y. Shao-Horn, *J. Phys. Chem. C*, 2013, **117**, 25926– 25932.
- 108 D. K. Bediako, Y. Surendranath and D. G. Nocera, J. Am. Chem. Soc., 2013, **135**, 3662–3674.
- 109 A. M. Ullman, C. N. Brodsky, N. Li, S.-L. Zheng and D. G. Nocera, J. Am. Chem. Soc., 2016, 138, 4229–4236.
- 110 Z. J. Xu, Sci. China Mater., 2020, 63, 3-7.
- 111 Y. Li, X. Du, J. Huang, C. Wu, Y. Sun, G. Zou, C. Yang and J. Xiong, *Small*, 2019, **15**, 1901980.
- 112 H. Li, Y. Chen, J. Ge, X. Liu, A. C. Fisher, M. P. Sherburne, J. W. Ager and Z. J. Xu, *JACS Au*, 2021, 1, 108–115.
- 113 H. Li, Y. Chen, J. Z. Y. Seow, C. Liu, A. C. Fisher, J. W. Ager and Z. J. Xu, *Small Sci.*, 2022, **2**, 2100048.
- 114 S. Song, H. Bao, X. Lin, X.-L. Du, J. Zhou, L. Zhang, N. Chen, J. Hu and J.-Q. Wang, *J. Energy Chem.*, 2020, 42, 5–10.
- 115 S. Ye, J. Wang, J. Hu, Z. Chen, L. Zheng, Y. Fu, Y. Lei, X. Ren, C. He and Q. Zhang, ACS Catal., 2021, 11, 6104–6112.
- 116 Q. Qin, H. Jang, Y. Wang, L. Zhang, Z. Li, M. G. Kim, S. Liu, X. Liu and J. Cho, *Adv. Energy Mater.*, 2021, **11**, 2003561.
- 117 Z. Wang, Y.-R. Zheng, I. Chorkendorff and J. K. Nørskov, ACS Energy Lett., 2020, 5, 2905–2908.
- 118 W. Zhu, F. Yao, K. Cheng, M. Zhao, C.-J. Yang, C.-L. Dong,
   Q. Hong, Q. Jiang, Z. Wang and H. Liang, *J. Am. Chem. Soc.*, 2023, 145, 17995–18006.
- 119 R. Li, B. Hu, T. Yu, H. Chen, Y. Wang and S. Song, *Adv. Sci.*, 2020, 7, 1902830.
- 120 K. Xiao, Y. Wang, P. Wu, L. Hou and Z. Q. Liu, Angew. Chem., 2023, 135, e202301408.
- 121 S. Liu, H. Tan, Y. C. Huang, Q. Zhang, H. Lin, L. Li, Z. Hu,
  W. H. Huang, C. W. Pao and J. F. Lee, *Adv. Mater.*, 2023,
  35, 2305659.
- 122 Z.-Y. Wu, F.-Y. Chen, B. Li, S.-W. Yu, Y. Z. Finfrock, D. M. Meira, Q.-Q. Yan, P. Zhu, M.-X. Chen and T.-W. Song, *Nat. Mater.*, 2023, 22, 100–108.
- 123 M. F. Lagadec and A. Grimaud, *Nat. Mater.*, 2020, **19**, 1140–1150.
- 124 H. B. Yang and B. Liu, Chem Catal., 2021, 1, 1365–1366.