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REVIEW

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Exploring the potential of cobalt hydroxide and its derivatives as a cost-effective and abundant alternative to noble metal electrocatalysts in oxygen evolution reactions: a review

Umair Shamraiz, (10)†*** Abdul Majeed,†** Bareera Raza,** Noor ul Ain and Amin Badshah^d

This review article provides a comprehensive analysis of the use of cobalt hydroxide and its derivatives as catalysts in the oxygen evolution reaction (OER), which is a crucial step in the water-splitting process for hydrogen production. The article covers the various synthetic methods used to prepare cobalt hydroxide and its derivatives, as well as their characterization techniques. The article also discusses the effect of different synthesis parameters on the performance of these materials as OER catalysts. The article goes on to evaluate the performance of cobalt hydroxide and its derivatives under various OER conditions,

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^aDepartment of Chemistry, Technical University of Darmstadt, 64287, Germany. E-mail: umair.shamraiz@tc1.tu-darmstadt.de; umairshamraiz@gmail.com

^bCollege of Chemistry and Environmental Engineering, Shenzhen University, Shenzhen 518060, China

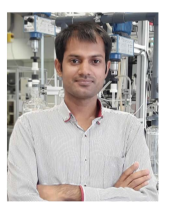
- Department of Chemistry, Rawalpindi Women University, Rawalpindi 46300, Puniab, Pakistan
- ^dDepartment of Chemistry, Quaid-i-Azam University, 45320 Islamabad, Pakistan † These authors contributed equally.



Umair Shamraiz

Umair Shamraiz, a distinguished researcher in multifunctional electro-catalysts for energy storage and conversion, spearheaded groundhas breaking advancements in the field. Specializing in metal oxide-based palladium/rGO electro-catalysts and α -CoOOH doped with transition metals, Umair has significantly contributed to the development of direct liquid fuel cells and facilitated oxygen evolution through water

splitting methodologies. Currently serving as a George Forster Postdoctoral Fellow at the esteemed Technical University of Darmstadt, Umair's pioneering work focuses on investigating novel avenues such as In₂S₃ and Ag-based thiospinels/tellurospinels for water splitting, pushing the boundaries of sustainable energy technologies. Driven by a vision for the future, Umair's aspirations include pioneering efforts in synthesizing electrolytes tailored for Mg-ion batteries and innovating catalysts that will revolutionize energy storage paradigms. With an unwavering commitment to scientific excellence, Umair Shamraiz continues to lead the charge in advancing the frontiers of electro-catalysis for a more sustainable energy landscape.



Abdul Majeed

Abdul Majeed, an esteemed scholar in Materials Science, earned his PhD from the prestigious CAS Institute in China in 2019. His doctoral research focused on the synthesis and exploration of diverse carbonbased materials specifically tailored for efficient water splitting processes, showcasing his expertise in sustainable energy Continuing solutions. academic journey, Abdul Majeed transitioned to a role as a Post-

doctoral Fellow at Shanghai Jiatong University. Here, his research delved into the electrodeposition techniques applied to a range of layered double hydroxide materials, advancing their potential applications in various energy-related fields. Presently serving as an Alexander Von Humboldt Postdoctoral Fellow at TU Darmstadt, Abdul Majeed's dedication to innovative research continues to shine. His significant contributions to the scientific community are evidenced by a portfolio of publications in esteemed journals such as ACS Applied Materials and Interfaces, as well as JMCA, solidifying his position as a distinguished authority in the field of materials science and energy conversion.

such as pH, temperature, and applied potential. The article also discusses the advantages and limitations of using these materials as OER catalysts, such as their stability, activity, and cost-effectiveness. The article compares the performance of cobalt hydroxide and its derivatives to other commonly used OER catalysts, such as iridium and ruthenium oxides.

1 Introduction

Electrocatalytic water splitting is a leading field in producing affordable hydrogen, which is currently the most important requirement for our environment as hydrogen is the greenest fuel.1 Hydrogen has properties such as the highest specific chemical energy density (142 MJ kg⁻¹), and the only byproduct of its use is water, unlike other systems, which release hazardous chemicals upon combustion.²⁻⁴ Additionally, we are facing environmental problems due to the excessive consumption of hydrocarbons, and known resources will be depleted by 2080.5,6 Therefore, it is crucial to move towards a hydrogen economy. However, hydrogen production is currently costly and energy-inefficient due to high temperature, pressure, complex technology, corrosive chemicals, and safety measures.7-9

Furthermore, it emits greenhouse gases directly into the air, similar to carbon and hydrocarbons. Steam reforming via hydrocarbons emits dangerous nitrogen, sulfur, carbon, and hydrogen, causing poor cell performance in fuel cells.7-15

The problem could be resolved by considering electrocatalytic water splitting, which produces a vast amount of energy from renewables. Additionally, it is the safest method for producing green and pure hydrogen. Water splitting is further divided into three different types, depending on the mode of energy used for catalysis: photocatalytic water splitting, 16-21 electrocatalytic water splitting,22-24 and photoelectrochemical water splitting.25 Compared with the electrocatalytic process, the other two are relatively slow in hydrogen production.

In electrocatalytic water splitting, three different types of electrolyzers are used, depending on their working principle.



Bareera Raza

As a dedicated Postdoctoral Fellow at Shenzhen University, Bareera Raza focuses on pioneering advancements rechargeable energy storage systems, particularly in the realm of non-aqueous electrolyte systems for zinc ion batteries. With a primary goal of developing safe, efficient, and highcurrent-capable electrolyte solutions, Bareera Raza conducts cutting-edge research aimed at pushing the boundaries of

battery technology. Bareera Raza is instrumental in the synthesis of energy-dense, high-voltage cathodes that prioritize compatibility within battery systems. Their work revolves around crafting cathode materials that optimize energy density while ensuring seamless integration with the overall battery architecture. Additionally, Bareera Raza employs a multidisciplinary approach, leveraging a diverse array of battery testing and characterization techniques. Through meticulous design, testing, and analysis, Bareera delves into the intricate chemistry of zinc ion batteries, unraveling insights critical for enhancing battery performance and durability. Driven by a passion for innovation, Bareera Raza stands at the forefront of research, contributing significantly to the development of next-generation rechargeable energy storage systems, shaping the landscape of sustainable energy technologies.



Noor ul Ain

Dr Noor ul Ain is an accomplished scholar, holding a distinguished PhD in Chemistry earned from Quaid-i-Azam University in April 2021. Her expertise lies at the intersection of inorganic chemistry, nanotechnology, and energy storage materials, particularly focusing on advancements in lithium-ion batteries. Currently serving as a lecturer (BS 18) at RWU since August 2022, Dr Noor ul Ain has established herself as a repu-

table researcher with contributions featured in esteemed international journals. Her academic prowess was further acknowledged through her selection for the IRSIP scholarship by HEC, which supported her groundbreaking research during her PhD at University College London. Dr Noor ul Ain's research interests encompass multifaceted environmental applications, specifically targeting the reduction of environmental carcinogens using nanomaterials. Additionally, her work extends to energy storage applications, where she specializes in synthesizing ternary metal oxides tailored as cathode materials for enhancing Li-ion batteries' performance. Moreover, her expertise encompasses the synthesis, characterization, and application of magnetic and non-magnetic copper sulfide nanoparticles, showcasing her versatility in nanomaterial research. Her research endeavors, ranging from environmental applications to advancements in energy storage solutions, are driven by a profound commitment to scientific exploration and innovation. Dr Noor ul Ain's contributions stand as a testament to her dedication to advancing knowledge and addressing critical challenges in the realms of chemistry, nanotechnology, and sustainable energy storage.

These (PEM) include proton exchange membrane electrolyzers, 26-29 anion exchange membrane-based alkaline electrolyzers,30-33 and high-temperature solid-oxide electrolyzers that operate on steam.34 Recently, water splitting in a neutral environment using buffered solutions has gained importance as it is more environmentally friendly and less corrosive. 23,34 However, these methods have some disadvantages, such as lower energy efficiency and the need for high cell potentials, and lower current density. On the other hand, many transition metal-based electrocatalysts have shown promising results for oxygen evolution reaction (OER) electrocatalysis under neutral conditions. 35-42

Among these electrocatalysts, those that can simultaneously catalyze both the anodic oxygen evolution reaction (OER) and cathodic hydrogen evolution reaction (HER) are considered superior. Many compounds, including metals and their composites, and carbon-based materials, have shown better activity for these reactions. However, the toxic and unstable nature of carbon-based materials hinders their industrial use for hydrogen production. 41,42 In contrast, metallic-based electrocatalysts are superior because they offer metallic surfaces enriched with electrons, a variety of oxidation states, and optimized metal-oxygen bond energies, which can facilitate the adsorption and desorption of intermediates. 43-46 So far, metals and their compounds such as Pt, 25,47,48 Ir, 47-50 Ru, 51-54 Ti, 55 V, 56-58 W, 59-62 Mo, 63-65 Mn, 66-68 Fe, 69-71 Co, 72-78 Ni, 79-83 Cu, 83,84 and Au85-87 have been successfully used for electrochemical water splitting. Moreover, only Pt is used as a metal, while the others are mostly used in the form of composites. The main purpose of Pt is to catalyze the HER because it can produce hydrogen at 0 V.87 However, it has recently been identified that

Prof. Dr Amin Badshah is a distinguished academician and researcher, having obtained his PhD in 1994 from Quaid-i-Azam University (QAU), Islamabad. With a prolific career in academia, he currently serves as the vice chancellor of the University of Buner and holds the esteemed positions of Ex-Professor Emeritus and Dean of the Faculty of Natural Sciences at QAU, Islamabad. A distinguished fellow of the Pakistan Academy of Sciences (PAS) and the Chemical Society of Pakistan, Dr Badshah's scholarly contributions extend to his role as an associate editor for the Journal of the Chemical Society of Pakistan, showcasing his commitment to advancing scientific discourse and research dissemination. His remarkable achievements in academia and research have been recognized through numerous accolades, including the Research Productivity Award from the Pakistan Council for Science and Technology spanning from 2001 to 2016. Dr Badshah's dedication to excellence has been further underscored by prestigious honors such as the Gold Medal from PAS, Best Teacher and Best Paper Awards from the Higher Education Commission of Pakistan, and the esteemed Tamgha-i-Imtiaz (TI) from the Government of Pakistan. Dr Badshah's illustrious career stands as a testament to his unwavering commitment to scholarly pursuits, academic leadership, and the advancement of scientific knowledge within Pakistan and beyond.

some phosphides and chalcogenides are capable of carrying out the HER in the same range as Pt. 88 On the other hand, Ru and Ir oxides are superior for the OER under acidic conditions.89,90 Unfortunately, these precious metals increase the cost of hydrogen production despite their efficiency. Research is underway to replace Pt with cheaper metals.89,90

Nevertheless, first-row transition metal electrocatalysts including hydroxides and layered double hydroxides of Co have been given preference due to their ease of preparation and better electrocatalytic performance compared to other types. 74,75,89,90 Cobalt-based hydroxides coupled with other metals have boosted oxygen evolution reactions. Therefore, a review article is needed to understand the catalysis phenomena of cobalt-based hydroxides and the factors that make them competitive with noble metals. Additionally, the challenges faced by Co-based hydroxides highlight the significance of the OER.91-94

Significance of electrocatalytic water splitting

Electrocatalytic water splitting, a non-spontaneous reaction, requires a 1.23 V thermodynamic potential, alongside kinetic hindrances necessitating an additional potential for the anode and cathode.95 The complex oxygen evolution process at the anode contrasts with the simpler hydrogen evolution at the cathode. The detailed mechanism of oxygen evolution involves different pathways and mechanisms like the adsorbate evolution mechanism (AEM) and lattice oxygen participation mechanism (LOM). Co-based hydroxides' catalytic capabilities are showcased, emphasizing their importance in oxygen evolution reactions.22,96

AEM

* +
$$H_2O(1) \rightarrow OH^* + H^+ + e^-$$

$$OH^* \rightarrow O^* + H^+ + e^-$$

$$O^* + H_2O(1) \rightarrow OOH^* + H^+ + e^-$$

$$OOH^* \rightarrow O_2 + H^+ + e^-$$

LOM

$$OH^* \to (Vo + OO^*) + H^+ + e^-$$

$$(Vo + OO^*) + H_2O(I) \to O_2 + (Vo + OH^*) + H^+ + e^-$$

$$(Vo + OH^*) + H_2O(I) \to (H^*_{o-site} + OH^*) + H^+ + e^-$$

$$(H^*_{o-site} + OH^*) \to OH^* + H^+ + e^-$$

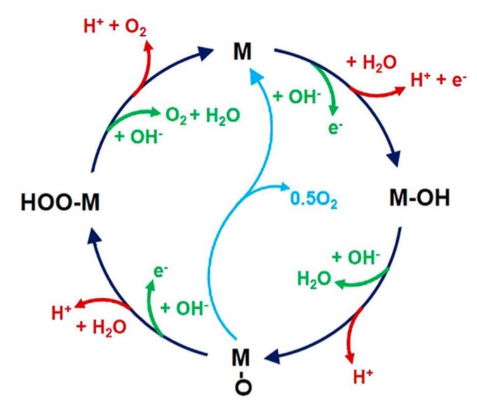


Fig. 1 Generalized mechanism of the OER in acidic (red path) and alkaline (green path) environments. This figure has been reproduced from ref. 97 with permission from American Chemical Society, copyright 2018.

In the context of these equations, 'Vo' denotes oxygen vacancies formed due to lattice oxygen involvement in catalysis. The OER mechanism, as depicted in Fig. 1 by Jin et al.,97 elucidates the necessity of four electrons and protons for oxygen evolution. This figure underscores that the OER demands catalysts with variable oxidation states and favorable metaloxygen bond energies, where cobalt hydroxide/layered double hydroxide stands as a noteworthy catalyst. It highlights the complexity of the OER compared to the HER, necessitating an additional potential for catalysis, typically within 250-500 mV, yet reaching up to 670 mV at an industrial scale. Efforts to diminish overpotential involve creating high surface area composites with varied catalyst loading. Modifying intrinsic activity via doping, particularly with metals like Fe, is another avenue that has been explored.12 Subsequent sections will emphasize the significance of cobalt hydroxide as an OER electrocatalyst under alkaline conditions.

Exploring the advantages of cobalt hydroxide in oxygen evolution reactions: electronic and structural properties

3.1 Polymorphs of cobalt hydroxide

Cobalt hydroxide exhibits distinct polymorphs, α-cobalt hydroxide and β -cobalt hydroxide. The β -form adopts a brucite-like structure with a Co(OH)₂ stoichiometry, featuring

a closed hexagonal arrangement of OH ions and Co(II) occupying alternating rows of octahedral sites. 98 Conversely, the α phase showcases a non-stoichiometric composition similar to hydrotalcite-like compounds. This phase includes positively charged $Co(OH)_{2-x}$ layers balanced by interlayer anions (e.g., Cl^- , CO_3^{2-} , NO^{3-} , S^{2-} , etc.), resulting in an extended interlayer spacing exceeding 7 Å, as opposed to the restricted 4.6 Å spacing in the β -form (Fig. 2).98,110

Notably, α-Co(OH)₂ demonstrates high specific capacity, making it a potential candidate for electrochemical supercapacitors and energy devices. 76,77,99,100 On the other hand, β-Co(OH)2's hexagonal layered structure presents an intriguing opportunity as a reactive template for thermoelectric cobaltite ceramics, such as NaCo₂O₄ and Ca_rCoO₂. 101-103 Density functional theory studies reveal the superior electrochemical activity of the α-phase, positioning it as a promising electrode material and a potential candidate for organic magnetic materials through the incorporation of organic anions. 104,105

The distinction between the α and β phases extends to their sites; while the β-phase hosts Co2+Oh sites exclusively, the αphase contains both Co²⁺Oh and Co²⁺Td sites. 106-108 This variation impacts their electrochemical behavior significantly, with the electroactive Co²⁺Td sites in the α-phase playing a crucial role. Synthesis of Co²⁺Td sites is considered vital for electrochemical processes, as indicated by UV-visible spectroscopy displaying distinct absorption bands for both sites in the α -phase, compared to a single absorption band in the β-phase (Fig. 3).¹⁰⁹

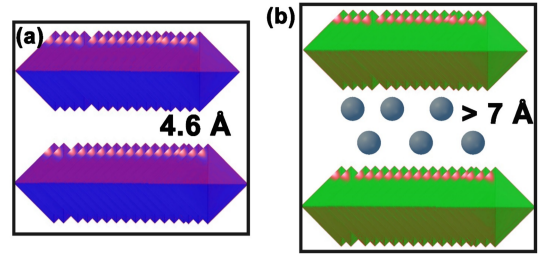


Fig. 2 Structure of (a) β -Co(OH)₂ and (b) α -Co(OH)₂

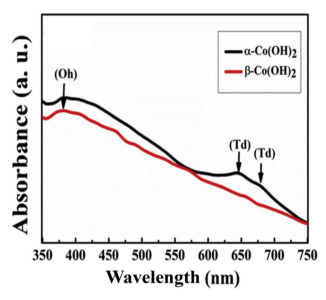


Fig. 3 UV-visible absorption spectrum of α - δ β -Co(OH)₂. This figure has been reproduced from ref. 109 with permission from Elsevier, copyright 2019

Cobalt hydroxide and cobalt oxyhydroxide

Cobalt hydroxide manifests in two polymorphs, distinguished by Co²⁺ ions occupying Co²⁺Oh and Co²⁺Td structural sites due to their distinct stoichiometries. Additionally, a structurally akin compound resembling α-Co(OH)2, known as cobalt oxyhydroxide, houses a network of Co²⁺ and Co³⁺ cations stabilized by interlayer anions in a hydrotalcite-like configuration. 117,118 This cobalt oxyhydroxide, represented by $[CoII_{(1-x)}CoIII_{(x)}]$ $OH_2[(A_x) \cdot nH_2O(A = NO^{3-}, Cl^-, CO_3^{2-} etc.)]$, accommodates both Co²⁺ and Co³⁺ on its layers. 117,119,120 The fabrication method and experimental conditions significantly influence the formation of these compounds. The α -Co(OH)₂, being a metastable state, gradually transforms into β-Co(OH)₂ or CoO(OH), a process heavily dictated by the OH-: Co2+ ratio. Fig. 4 illustrates this

transformation mechanism. Under specific OH⁻: Co²⁺ ratios, the polymorphs' conversion from α-Co(OH)₂ to β-Co(OH)₂ occurs gradually, while higher ratios prompt immediate transformation. Elevated temperatures expedite the conversion of α - $Co(OH)_2$ to β - $Co(OH)_2$, evident even at lower OH^- : Co^{2+} ratios, indicating a faster transformation at higher temperatures.

Investigating the use of cobalt hydroxide as an OER electrocatalyst in water splitting

4.1 β-Cobalt hydroxide (β-Co(OH)₂)

In our general discussion, the review starts with β -Co(OH)₂, as it is the most stable polymorph and is commonly produced by researchers, either intentionally or unintentionally.

4.1.1 β-Cobalt hydroxide nanoplates as OER electrocatalysts. Koza et al. 111 successfully prepared β-Co(OH)2 by the electrochemical reduction of tris(ethylenediamine)cobalt(III) at room temperature. They confirmed the formation of pristine β-Co(OH)2 via PXRD analysis, where all the peaks were indexed to brucite-like Mg(OH)₂. It is understood that β -Co(OH)₂ is stoichiometric and crystalline in nature, which can be seen in the SEM analysis; conical-shaped well-crystallized structures were identified. Similarly, the XPS analysis shows five deconvoluted peaks for Co, which match well with Co(OH)2, and oxygen has two binding modes for OH and Co-O bonds illustrated in Fig. 5. Oxygen intermediates' binding modes, OH and Co-O, are pivotal in the OER, dictating their efficiency and kinetics. Moreover, OH assists adsorption, stabilizing intermediates, while Co-O binding affects surface reactivity; optimal binding balances stability and turnover, crucial for efficient OER kinetics.

For electrochemical purposes, they deposited the material on a Ti electrode and LSV curves were measured for both Co₃O₄ and Co(OH)2 for comparative purposes. Both electrodes showed a similar profile, and Co₃O₄ shows better activity per geometric

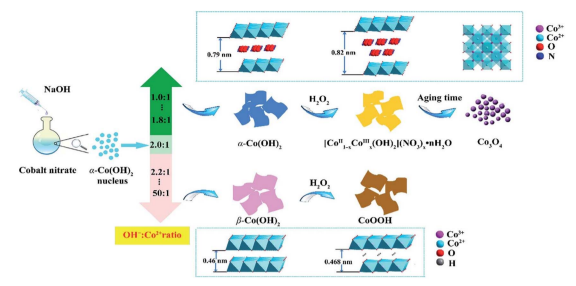


Fig. 4 Illustration of the formation of Co₃O₄ and CoOOH crystals. This figure has been reproduced from ref. 110 with permission from Royal Society of Chemistry, copyright 2011.

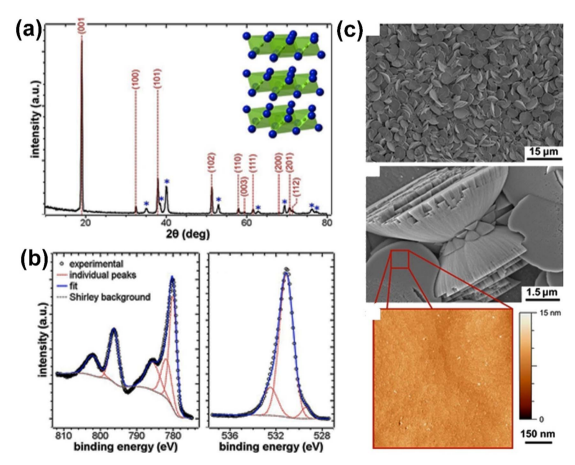


Fig. 5 The results of the XRD, XPS, and morphology analysis of electrodeposited Co(OH)₂. (a) Shows the XRD pattern of the Co(OH)₂, with the reflections of Co(OH)₂ indicated in red (JCPDS no. 30-0443) and the blue asterisks representing the Ti substrate. The inset of the panel shows the structure of the Co(OH)₂. (b) Shows the XPS spectra of the electrodeposited Co(OH)₂, with the Co 2p and O 1s binding energy ranges represented. (c) Shows the morphology of the $Co(OH)_2$ electrodeposited at a charge density of 0.6 C cm^{-2} , including SEM images of the film and an AFM image of a flat disk surface. This figure has been reproduced from ref. 111 with permission from American Chemical Society, copyright 2013.

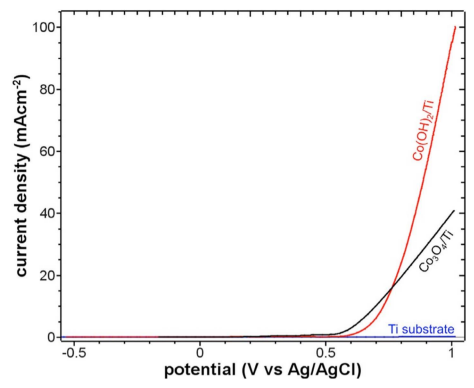


Fig. 6 LSVs measured at 1 mV s⁻¹ scan rate in 1 M KOH at the electrodeposited Co(OH)₂ (red), crystalline Co₃O₄ (black), and uncoated Ti substrate (blue). This figure has been reproduced from ref. 111 with permission from American Chemical Society, copyright 2013.

surface area (Fig. 6). However, the activity was undoubtedly increased for Co(OH)2 when optimized to the active surface area. This enhanced activity was attributed to the roughness factor possessed by the Co(OH)₂ in comparison with the smooth Co₃O₄. But in our view, it could be mere speculation, as no experimental evidence is provided to prove the conclusion, and the SEM shows very smooth cones of Co(OH)2.

Zhan et al.¹¹² synthesized β-Co(OH)₂ hexagonal nanoplates via the well-known hydrothermal method and exploited them for the OER in an alkaline medium. The phase purity was confirmed via PXRD analysis, showing only the presence of peaks for β -Co(OH)₂. Furthermore, they applied heat treatment on the synthesized β -Co(OH)₂ to make CoO and Co₃O₄, which was confirmed by PXRD analysis and demonstrated that under the corrosive heat treatment, the OH⁻ molecules leave the surface leaving behind pores to increase the surface area, confirmed by BET analysis. The TEM analysis showed the presence of uniform hexagonal plates having an approximate thickness of 14 nm with an aspect ratio of 5.7. The LSV curves show that the OER activity is much better with Co(OH)2 in comparison with CoO and Co₃O₄. The Co(OH)₂ attained 10 mA cm⁻² OER activity at an overpotential of 450 mV, while CoO and Co₃O₄ required 460 and 510 mV overpotentials, respectively. Moreover, after 400 cycles, the activity was lost, and poststability characterization confirmed the conversion of Co(OH)₂ to Co₃O₄ under prolonged alkaline conditions. Post-stability analysis revealed Co(OH)2 oxidation to Co3O4 under extended anodic exposure: XRD indicated the presence shift from Co(OH)₂ to Co₃O₄, supported by XPS measurements showing Co

 $2p_{1/2}$ and Co $2p_{3/2}$ binding energy shifting from 798.5/783.0 eV to 794.8/779.8 eV, respectively. This oxidation mechanism elucidates Co(OH)₂ degradation under prolonged anodic potential, necessitating strategies to enhance its stability for sustained performance. The corresponding Tafel analysis shows faster reaction kinetics of the OER on the surface of Co(OH)2 as compared with the CoO and Co3O4 owing to the Tafel slope values of 60 mV dec⁻¹, 65 mV dec⁻¹ and 75 mV dec⁻¹, respectively. Using hydroxides over oxides simplifies achieving excellent bifunctional activity; Co(OH)2 showcases superior performance due to optimal Co-OH bond strength, facilitating efficient oxygen electrocatalysis by bypassing certain kinetic steps. This study proposes $Co(OH)_2$ as a more costeffective and potent alternative to complex oxides like MnCoFeO₄, offering competitive ORR and OER capabilities, positioning it favorably against precious metal-based bifunctional catalysts.

4.1.2 β-Cobalt hydroxide nanosheets as OER electrocatalysts. β-Co(OH)₂ nanosheets were synthesized by Tong et al.113 They designed β-Co(OH)2 under three different experimental conditions and proposed that the crystallinity of the samples increased with the increase in temperature and reaction time, as supported by PXRD analysis. However, the thickness of the sheets increased from 3.9 nm for a reaction time of 12 hours at 120 °C to 41.7 nm for a reaction time of 16 hours at 160 °C. The structural analysis of the acquired Co(OH)₂ specimens underwent examination through SEM, TEM, and AFM techniques. Displayed in Fig. 7b is the SEM image that portrays a characteristic expansive view of the Co(OH)2 product,

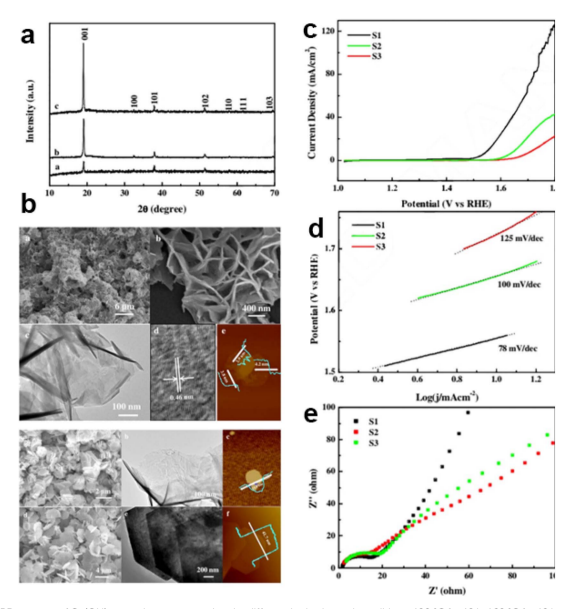


Fig. 7 (a) XRD patterns of Co(OH)₂ nanosheets prepared under different hydrothermal conditions: 120 °C for 12 h, 160 °C for 12 h and 160 °C for 16 h. (b) SEM, TEM and AFM images of Co(OH)₂ nanosheets prepared under hydrothermal conditions at 160 °C for 12 h; SEM, TEM and AFM images of Co(OH)₂ nanosheets prepared under hydrothermal conditions at 160 °C for 16 h. (c) Polarization curves, (d) Tafel plots and (e) electrochemical impedance spectra. This figure has been reproduced from ref. 113 with permission from Elsevier, copyright 2017.

revealing a consistent and finely structured 2D form. Fig. 7b, with increased magnification, further authenticates the uniformity and micrometer-scale dimensions of the structure, exhibiting a thickness below 10 nm. The TEM depiction in Fig. 7b presents an abundance of mesh-like nanosheets with upturned edges due to heightened surface tension. The highresolution TEM in Fig. 7b reveals distinct lattice fringes indicative of well-crystallized phases, showcasing a spacing of 4.6 Å, corresponding to the (001) planes of hexagonal Co(OH)2. This confirms the single-crystalline nature of the Co(OH)₂ nanosheets.

AFM analysis was carried out to determine the thickness of the 2D Co(OH)₂ nanosheets, and Fig. 7b displays tapping-mode images, revealing an average vertical height of approximately

4.0 nm. This direct confirmation underscores the ultrathin attribute of the acquired Co(OH)2 nanosheets. The Co(OH)2, termed as ultrathin nanosheets, produced a current density of 10 mA cm⁻² at only 327 mV overpotential, and the kinetics were relatively faster with a Tafel slope of 78 mV dec^{-1} for the sample with a thickness of 3.9 nm. The authors attributed this enhanced activity to the fine, thin nanosheets.

The ECSA calculations suggested the largest surface area for the sample synthesized at lower temperatures, which is the main contributing factor towards its better activity. The increase in thickness results in an increase in both onset potential and overpotential at 10 mA cm⁻², illustrating that morphology and size have a pronounced effect on the intrinsic activity of the electrocatalysts. This effect is also seen in the

the nanosheets, but the author did not draw a conclusive statement for this enhancement. In conclusion, the better activity could be due to the NF electrode used, having high surface and porosity results which have a pronounced effect not only on the activity but also on the stability of the material as well. Ghanem et al. 118 used a liquid crystal template to fabricate mesoporous cobalt hydroxide. Due to the oxidation peak of cobalt, the authors reported 25 mA cm⁻² at 1.52 V, which is 130 mV lower in comparison with bulk cobalt hydroxide. Nevertheless, it shows that the overall overpotential should be lower, between 250 and 270 mV at a current density of 10 mA cm⁻². Moreover, they performed the OER in 2.0 M KOH as well, which displayed even better activity, and a current density of 20 mA cm⁻² was achieved at 1.50 V. It is obvious that the enhanced activity was due to the increased electrochemically active surface area because of the porous nature of the material.

Yang et al. 119 explored Co-OOH bond elongation in Co(OH)2 nanosheets via crystal distortion for enhanced OER. By incorporating Ga (gallium) into the Co(OH), there's a unique alteration in the crystal lattice that stretches out the Co-OOH bonds specifically on the 120 facets, generating active sites. Despite Ga's inclusion showing no distortion in the parent structure, PXRD revealed peak shifts, enhancing the OER due to longer metal-oxygen bond lengths. Catalysts exhibited exceptional OER at 265 mV overpotential for 10 mA cm⁻² with improved conductivity. Zhang et al. 120 designed self-supported hexagonal β-cobalt hydroxide via a painting-alloying-dealloying method, attaining a low 332 mV overpotential for 10 mA cm⁻² in the OER without a binder. The ultrathin structure and in situ growth on Ga-coated Co plates enhanced electronic conductivity and electrolyte movement. Dileep et al. 121 exfoliated β-Co(OH)2 to smaller sheets, achieving 390 mV overpotential for 10 mA cm⁻². The group pf Coleman¹²² used liquid exfoliated Co(OH)₂, obtaining 440 mV overpotential. Nanosheet optimizations in length, edges, and loading concentrations improved activity. Anchoring on carbon nanotubes reduced overpotential to 295 mV, demonstrating superior OER.

4.1.3 β-Cobalt hydroxide in other morphologies as OER electrocatalysts. Although LDHs mostly exist in the form of sheets, other morphologies do exist and there is sufficient data available that should be included in the review. Zhang et al. 123 fabricated sulfur Co(OH)2 amorphous nanoflakes on nickel foam and used them as an efficient OER electrocatalyst. They demonstrated that optimized sulfur engineering along with the nanoflake morphology is the deciding factor for enhanced activity. The overpotential was merely 283 mV for a current density of 100 mA cm $^{-2}$. Nevertheless, they further measured a higher current density of 1000 mA cm⁻², which required an overpotential of 365 mV. The incorporation of sulfur improved the Co³⁺/Co²⁺ ratio, which engendered the *OOH intermediates. The speculations were further supported by DFT calculations revealing that the sulfur withdraws electronic density from the Co atoms, hence reducing the free energy of the *OOH intermediates, leading to an increase in intrinsic activity. However, the authors did not relate the activity to the amorphous and nanoflake nature of Co(OH)2. Hence, we can conclude that future researchers who are interested in OER electrocatalysis

highest TOF of the sample with the lowest thickness among the assembly of electrocatalysts tested. Moreover, the authors claimed that at elevated temperatures, hexagonal nanoplates are a favored morphology. The nanoplates have a thickness much higher than those of the other two samples in the series and have inferior activity. These results are consistent with the results discussed above reported by Zhan et al.112 Hence, nanosheets with an amorphous nature are better electrocatalysts for the OER than well-defined hexagonal nanoplates.114 The smooth surface does not offer effective participation of the underlying atoms during electrocatalysis, while a rough surface is advantageous as it allows more atoms to contribute to the process, resulting in a better activity. These results are also supported by the synthesis of amorphous Co(OH)₂ by the group of P. O'Mullane. 115 They reported enhanced OER activity when they subjected the amorphous Co(OH)2 as an electrocatalyst requiring an overpotential of only 360 mV to generate a current density of 10 mA cm⁻² with a Tafel slope of 58 mV dec⁻¹. These two results are consistent with the enhancement of electrochemically active surface area due to the rough and amorphous nature of the electrocatalysts. However, Tong et al. 113 proposed that no transformation of the parent β-Co(OH)₂ occurred during the OER. Moreover, they have suggested that the Co(OH)2 is converted into Co₃O₄ during the OER. Furthermore, to support their results and to make a better scientific contribution, they have tested their catalysts for the OER at different electrodes. They demonstrated that the Au electrode exhibits superior OER electrocatalysis as compared with the Pd, Cu, and GC electrodes, and the order of activity was found to be Au > Pd > Cu.

Moreover, they supported their results with the role of electronegativity of the electrodes subjected to the OER. Among these electrodes, Au is more electronegative and will definitely facilitate the oxidation of Co(II) to the active Co(III) and then to Co(IV) during the OER as compared to the less electronegative electrodes in the series. These results are also supported by DFT calculations which demonstrate that Au positively shifts the dband center of Co by 0.74 eV compared to pure Co. The elevated d-band center favors the oxidation of Co. 116 Similarly, Babar et al.117 fabricated Co(OH)2 nanosheets directly on Ni foam by electrodeposition and subjected it to heat treatment up to 400 °C in air. The better activity of Co(OH)2/NF over Co3O4/NF was attributed to the enhanced surface area, availability of more active sites, and better charge transport ability. These results are in line with the results discussed so far in this review. The annealing of the electrodeposited Co(OH)2/NF catalysts has two pronounced effects: there is a transformation of Co(OH)2 to Co₃O₄ and also the crystallinity of the sample is increased. This is the main reason why Co(OH)₂ is a potent candidate for the OER; the amorphous nature is the driving force behind this superiority. Along with the amorphous nature, it was also identified that annealing has reduced the interlayer spacing between the layers owing to the aggregation, which results in the reduction of the active surface area. These results suggested that the activity enhancement was due to the amorphous and porous nature of Co(OH)2, both of which were changed upon annealing and resulted in the loss of activity. The OER activity reported here is much better than the reports so far dealing with

should consider the factor of electron-withdrawing groups in the material because the OER is highly dependent on how electron-deficient the metal center is.124

Zhang et al. 125 synthesized needle-like 3D carbonateincorporating Co(OH)₂ superstructures for the OER. The synthesis was a simple hydrothermal treatment using CoCl₂-·6H₂O as a source of cobalt, urea as a precipitating agent, and sodium glycolate as a surfactant. The reaction was carried out at 150 °C. The 3D structure is formed by the nucleation of the 1D needle-like structures. The authors did not emphasize the factors responsible for the formation of these unique 3D structures. In conclusion, the most probable factor could be the use of the sodium glycolate surfactant, which could be responsible for the formation of needle-like 3D structures. The 3D superstructures delivered a current density of 10 mA cm⁻² at the expense of merely 240 mV overpotential. The authors drew a few conclusive points depending on the reaction conditions. The apparent size of the CCHS appeared to be in micrometers, but they are constructed from the nanoneedles having a fine nanometer size range. Nevertheless, the CCHS has an influence on the transport of the electrolyte onto the active sites owing to good electrolyte-surface compatibility. Furthermore, the samples synthesized at 150 °C for 15 h and 120 °C for 12 h agglomerated quickly, and the structures were too thick to transport electrolytes effectively. Moreover, the samples synthesized at 150 °C for 6 h and 180 °C for 12 h appeared to be too loose for ion and electron transport.

Darbandi et al. 126 reported hexagonal mesoporous rings of β-Co(OH)₂ by a chemical precipitation method. The ring-like structure is formed with the H₂O₂-assisted method, while the rest of the procedure is similar to the reports discussed above. The rings appeared as aggregates of small fine nanoparticles.

Dhawale et al. 127 adopted a microwave-assisted synthesis for the fabrication of Co(OH)2 nanorods and later on, their thermal conversion into porous Co₃O₄ nanorods. They used CTAB as a surfactant during the synthesis of Co(OH)2. CTAB is a very powerful surfactant for making the nanorod-like morphology of nanomaterials. They tested the OER in 0.1 M KOH solution. The sample calcined at 600 °C showed better performance as compared to the other samples in the list. The authors claimed that the enhanced performance was due to the increase in active sites at a higher temperature. Liu et al.128 fabricated Co(OH)2 nanocages and obtained a current density of 10 mA cm⁻² at an overpotential of 280 mV.

4.2 α -Cobalt hydroxide (α -Co(OH)₂)

In the subsequent sections the discussion will be focused on the impact of different morphologies of α-Co(OH)2 on the oxygen evolution reactions.

4.2.1 α-Cobalt hydroxide nanosheets as OER electrocatalysts. Huang et al. 129 provided a detailed analysis of the electrocatalytic activity of various materials toward the OER using several techniques. LSV was used to obtain polarization curves of the α -Co(OH)₂-GNS-x and commercial RuO₂ catalysts. The LSV results show that the α-Co(OH)2-GNS-3 catalyst has a notably enhanced electrochemical activity compared to the

other samples, as evidenced by its lower onset potential (E_{onset}) of 1.42 V, which is the potential required to achieve a current density of 0.1 mA cm⁻². Additionally, the α -Co(OH)₂-GNS-3 catalyst has a lower overpotential of 259 mV at 10 mA cm⁻², indicating that it requires less energy to drive the reaction. It is found that the α-Co(OH)₂-GNS-3 has the lowest Tafel slope, 85.9 mV dec⁻¹, which further indicates that it has the most favorable intrinsic reaction kinetics. Electrochemical impedance spectroscopy (EIS) analysis is employed to interpret the OER capability of the α -Co(OH)₂-GNS-x from a dynamic point of view. The Nyquist plots suggest that the α-Co(OH)2-GNS-3 has the lowest impedance value (2.1 Ω) and a rapid proton-coupled electron transfer behavior compared to other catalysts. Cyclic voltammograms are used to study the double-layer capacitance $(C_{\rm dl})$ of the catalyst at different scan rates between 20 and 100 mV s⁻¹, which is used to calculate the electrochemically active surface area (ECSA) of the catalysts. The results show that the α -Co(OH)₂-GNS-3 has the highest C_{dl} (59 mF cm⁻²) compared to the other catalysts, which corresponds to the highest ECSA value of 1475. Overall, the authors present a comprehensive analysis of the electrocatalytic activity of the α - $Co(OH)_2$ -GNS-x and commercial RuO₂ catalysts towards the OER using several techniques, and the results indicate that the Co(OH)2-GNS-3 catalyst has superior electrocatalytic activity compared to the other samples.

Bu et al.130 studied the OER activity of Co₃(PO₄)₂ (CP), Co(OH)₂ (CH), Co₅(PO₄)₄OH₂ (CPH), and commercial RuO₂. The study uses linear sweep voltammetry and Tafel plots to evaluate the performance of the catalysts. The LSV results show that CP and CH have relatively high overpotentials (370 and 360 mV, respectively) at a current density of 10 mA cm⁻², whereas CPH and RuO2 have significantly lower overpotentials (254 and 256 mV, respectively) at the same current density illustrated in Fig. 8a. The Tafel plots give further insight into the reaction kinetics. The Tafel plots show that CPH has a smaller Tafel slope (57 mV dec^{-1}) than CH (210 mV dec^{-1}) and CP (246 mV dec^{-1}) in the range of high overpotentials, indicating that CPH is more active for the OER as illustrated in Fig. 8b. This can be attributed to the presence of phosphorus and hydrogen in CPH, which helps to create more active sites and improve the catalytic activity. Analysis describes the reaction mechanism for the OER in Co-based catalysts such as CP, CH, and CPH, which involves bonding interactions within intermediates (M-OH, M-O, and M-OOH) followed by O-O formation. The relatively large Tafel slopes of CH and CP suggest that the rate-limiting step of the OER process is in the first step of this mechanism, where one electron is transferred, which is consistent with previous reports. The small Tafel slope of CPH, on the other hand, suggests that the rate-limiting step is different, indicating that CPH is highly active for the OER. The electrochemically active surface area (ECSA) plays an important role in electrocatalytic reactions. CPH has $C_{\rm dl}$ of 14 mF cm⁻², which is nearly 3.5 times and 2.3 times larger than those of CP (4 mF cm⁻²) and CH (6 mF cm⁻²), respectively shown in Fig. 8c. This suggests that CPH has more electrochemical active sites and a larger surface area than CP and CH, which improves its electrocatalytic performance. Additionally, the EIS plots revealed that the ionic and ohmic

resistance of CPH is smaller than that of CP and CH, indicating that CPH has better electrical conductivity, which contributes to its excellent OER performance demonstrated in Fig. 8d. Moreover, the ECSA normalized current densities also revealed that the CPH has a lower onset potential as compared with the other samples in the series shown in Fig. 8e. The CPH was further evaluated for a long-term stability test at different current densities such as 10 mA cm⁻² and 30 mA cm⁻², and it showed excellent stability under alkaline conditions illustrated in

Cui et al. 131 investigated the electrocatalytic activity of α-Co(OH)₂ nanosheets towards the OER in a 0.1 M KOH solution. The nanosheets were prepared using a MOF-mediated method and were characterized in terms of surface area, electrocatalytic activity, and stability. For comparison, they also investigated samples prepared with urea (Co(OH)2-urea) and with a 4 hour solvothermal treatment (Co(OH)₂-4 h) using the same mass loading (0.25 mg cm⁻²). The results showed that the prepared α -Co(OH)2 nanosheets (Co(OH)2-12 h) had the lowest overpotential of 340 mV at a current density of 10 mA cm⁻², while the Co(OH)₂-urea had the highest overpotential of 480 mV. The study also found that the Tafel slope of Co(OH)2-12 h was significantly lower than that of Co(OH)2-urea, indicating more favorable reaction kinetics. Furthermore, it was found that the surface area of the samples correlated with their OER activity, with higher surface area samples exhibiting better activity. The study suggests that the well-dispersed structure and highly

exposed surface area of the prepared α-Co(OH)2 nanosheets were responsible for their superior OER activity compared to the other samples. The results were supported by measurements of ECSA and EIS. Moreover, the prepared α -Co(OH)₂ nanosheets had high stability under OER conditions.

Jash et al. 132 investigated the electrocatalytic activity of different materials for the OER in a 1 M NaOH electrolyte solution at pH 13.9. They used CV to measure the activity of the materials and compared them. The exfoliated single layer (SL-Co(OH)₂) required the lowest overpotential (350 mV) to achieve a current density of 10 mA cm⁻², which is significantly lower than those of α -o(OH)₂ (380 mV) and β -Co(OH)₂ (437 mV). The Tafel slope also decreased from 85 to 57 mV dec⁻¹ from β-Co(OH)₂ to SL-Co(OH)₂, further indicating that SL-Co(OH)₂ is the best catalyst among this series of catalysts. SL-Co(OH)2 had a remarkable mass activity of 153.8 A g^{-1} and an intrinsic activity parameter TOF of 0.146 s⁻¹, which were 3.8 and 16.2 times higher for SL-Co(OH)₂ than those of α -Co(OH)₂ and β -Co(OH)₂ respectively. These results suggest that SL-Co(OH)₂ is a more active catalyst for the OER. The improved conductivity and exposed metal ions of SL-Co(OH)2 are the reason behind this high catalytic activity, as they provide more active sites to tune the band gap states and support the electron transfer during the OER process.

In an interesting study Huang et al. 133 used XAFS to investigate the reaction mechanism of the OER on α -Co(OH)₂. In this study they used LSV to measure the OER activity of an α-

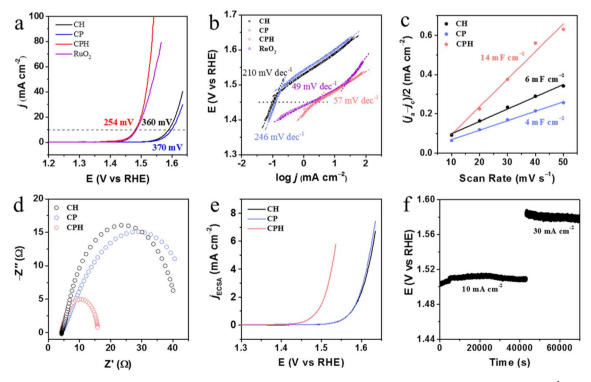


Fig. 8 (a) OER polarization curves of CP, CH, and CPH deposited onto glassy carbon electrodes. The sweep rate is 5 mV s⁻¹ in 1 M KOH. (b) Corresponding Tafel slope plots. (c) ECSA evaluation from the plots of current density at 0.15 V vs. RHE. (d) EIS spectra of all samples at an overpotential of 350 mV. (e) Specific current densities of all samples obtained via normalizing the geometric current densities to the ECSA. (f) Stability test of CPH at the current density of 10 mA cm⁻² and then 30 mA cm⁻². This figure has been reproduced from ref. 130 with permission from American Chemical Society, copyright 2019.

Co(OH)₂ nanosheet in 1 M KOH electrolyte. The α-Co(OH)₂ nanosheet had a notably small overpotential of 320 mV at a current density of 10 mA cm⁻² for the OER, which is lower than that of the state-of-the-art IrO2 electrocatalyst (overpotential of 340 mV at 10 mA cm⁻²). The α -Co(OH)₂ nanosheet had excellent durability under alkaline conditions and it exhibited good electrochemical stability. To understand the relationship between the structure and activity of the α -Co(OH)₂ nanosheet, the researchers used X-ray absorption fine structure (XAFS) measurement to characterize its oxidation state and structure before and after the OER. The results of XAFS measurement suggested that the α-Co(OH)₂ nanosheet undergoes a phase change from α-Co(OH)₂ to γ-CoOOH during the OER procedure, with an increased oxidation state and proportion of Co sites that contain a formal oxidation state of Co(IV). The results of the XRD pattern and EXAFS data also confirm this phase change. This suggests that the excellent catalytic activity of the α -Co(OH)₂ nanosheet originated from the formation of γ -CoOOH, which is facilitated by the presence of O vacancies in the surface Co octahedron during the OER, thus leading to the formation of hydroperoxy (OOH) species.

The detailed analysis of the study done by Jiang et al. 134 showed that the electrocatalytic activity of PI/CNT-Co(OH)2 films was investigated in 1 M KOH using a standard threeelectrode system. The Co(OH)2 2 min film showed an lower onset potential at around 1.50 V vs. RHE with a rapid increase of OER current. In comparison, the $Co(OH)_2$ products at 0.5, 1, 3, 4, and 5 min had higher onset potentials, and their OER current dropped below that of the Co(OH)2 2 min film. The Co(OH)2 2 min film generated a current density of 10 mA cm⁻² at an overpotential of 317 mV, whereas the Co(OH)₂ products at other deposition times exhibited more positive overpotentials. The prior peak of Co(OH)₂ around 1.05 V was assigned to the Co(II)/ Co(III or IV) redox process. In the electrochemical process, Co(OH)2 was first oxidized to CoOOH, and then the CoOOH was further oxidized to CoO2. Both CoOOH and CoO2 can catalyze the OER, with CoO₂ being more efficient for the OER. The electrochemical process of Co(OH)2 at the PI/CNT film can be described by the following reactions:

$$Co(OH)_2 + OH \rightarrow CoOOH + H_2O + e$$

 $CoOOH + OH \rightarrow CoO + H_2O + e$

The Co(OH)₂ 2 min film showed the lowest overpotential and the fastest increase of OER current among the products, indicating that the PI/CNT-Co(OH)2 2 min film exhibited better electrocatalytic activity for the OER, mainly determined by the catalytically active sites of Co(OH)2 nanosheets. With the deposition time increasing from 0.5 to 2 min, an increasing number of Co(OH)2 nanosheets were formed on the PI/CNT film, providing a large amount of active sites for the OER. However, further extension of the reaction time (3-5 min) resulted in the aggregation of nanosheets, decreasing the active site density of Co(OH)2 and weakening the OER electrocatalytic activity of Co(OH)₂. To obtain further insights into the OER

activity, Tafel plots of PI/CNT-Co(OH)2 films at different deposition times were studied. The Co(OH)2 2 min film showed a small Tafel slope of 49 mV dec⁻¹, which is close to the Tafel slopes of well-established OER catalysts Ir/C (40 mV dec⁻¹) and IrO_2 (49 mV dec⁻¹). The Co(OH)₂ products at 0.5, 1, 3, 4, and 5 min displayed relatively higher Tafel slopes of 87, 70, 57, 65, and 79 mV dec^{-1} .

Liang et al. 135 revealed that the ultrathin α-Co(OH)2 nanosheets have a high electrocatalytic activity for the OER due to their unique structure and composition. The ultrathin α-Co(OH)₂ nanosheets have a thickness of around 50 nm and a hexagonal plate-like structure, which allows for a large surface area and efficient charge transfer. Additionally, the Tafel slope of the α -Co(OH)₂ nanosheets was 64.9 mV dec⁻¹, which is much smaller than that of hexagonal α-Co(OH)2 plates (81.2 mV dec⁻¹) and commercial RuO₂ (78.7 mV dec⁻¹), indicating superior reaction kinetics in terms of OER. Furthermore, pHdependent studies of the α -Co(OH)₂ nanosheets revealed that the current density increases with increasing pH values, which indicates that the Co species in the nanosheets are oxidized easily at high concentrations of OH-. This suggests that the ultrathin α-Co(OH)₂ nanosheets may have potential applications in pH-sensitive electrocatalytic systems. α-Co(OH)₂ nanosheets had good durability, as demonstrated by the CPE test. After galvanostatic conditioning at 1.65 V for about 25 h, the current density was still maintained at around 11 mA cm⁻², indicating good stability of the α-Co(OH)₂ nanosheets. Overall, the study suggests that the ultrathin α -Co(OH)₂ nanosheets have a high electrocatalytic activity for the OER, which is attributed to their unique structure, composition, and pHdependent behavior. Ultrathin α-Co(OH)2 nanosheets have good durability and stability. These findings may have potential implications for the development of efficient and durable electrocatalysts for water splitting and other electrochemical applications.

Mao et al.136 investigated the electrocatalytic activity of a modified electrode composed of α -Co(OH)₂, polypyrrole (PPy), and graphene oxide (GO) for the OER in an alkaline solution. LSV was used to measure the OER activity of the modified electrode and compare it to other modified electrodes. The results showed that the PPy modified α-Co(OH)₂ electrode had a much lower onset potential for the OER than other modified electrodes (262 mV compared to 523 mV for the GO-modified GCE), indicating that the addition of PPy significantly improves the activity of the electrocatalyst. Moreover, the electrical conductivity of the as-synthesized pristine PPy powders is $0.6 \, \mathrm{S} \, \mathrm{cm}^{-1}$, while the conductivity of pure GO and PPy/GO is $0.5 \, \mathrm{G}$ \times 10⁻⁴ and 1.2 \times 10⁻⁴ S cm⁻¹, respectively. This information helps to explain why the addition of PPy improves the conductivity of the modified electrode, allowing for more efficient electron transfer and promoting the OER in the positive direction. Furthermore, they determined that the double-layer capacitance of the α-Co(OH)₂/PPy/GO modified GCE is 15.6 mF cm⁻², which is higher than that of the α -Co(OH)₂ modified GCE (3.53 mF cm⁻²) and α -Co(OH)₂/GO modified GCE (11.0 mF cm⁻²). This further supports the conclusion that the modified electrode has a higher electrochemically active surface area due

to the addition of PPy. Additionally, the Tafel slope of the α -Co(OH)₂/PPy/GO modified GCE is 74.5 mV dec⁻¹, much lower than those of GO (387 mV dec⁻¹), PPy/GO (168 mV dec⁻¹), α - $Co(OH)_2$ (94.8 mV dec⁻¹), and α -Co(OH)₂/GO (84.3 mV dec⁻¹).

Wang et al.137 reported the OER activity of a composite material made of graphene nanoribbons (GNRs) and α-Co(OH)₂. The study evaluated the OER activity of the composite material using a three-electrode system in a 1 M KOH solution. The results show that the GNRs/Co(OH)₂ composite had the lowest onset potential of 1.46 V (versus RHE) and an overpotential of 280 mV for reaching a current density of 10 mA cm⁻². This is in contrast to the control samples, with a-Co(OH)₂ alone showing a larger overpotential of 350 mV at 10 mA cm⁻², and GNRs exhibiting a sluggish polarization due to their inertness for the OER process. A combination of GNRs and Co(OH)₂ leads to a synergetic effect, with the GNRs providing good electron conductivity and the Co(OH)₂ being an active OER catalyst. The study also suggests that the morphology and state of the Co(OH)2 nanoparticles formed on the GNRs play an important role in their OER activity. Specifically, the thin and well-dispersed nanoparticles on the GNRs lead to more exposed surface active sites and better contact with the substrate. This is supported by TEM analysis, which showed that thin and entangled Co(OH)2 nanoparticles are formed on the GNR substrate, while irregular and large Co(OH)2 particles are produced when CNTs are used as the substrate. Furthermore, GNRs/Co(OH)₂ displayed a lower slope of 66 mV dec⁻¹, corresponding to the fastest kinetics, compared to CNT/Co(OH)₂ (81 mV dec⁻¹), α -Co(OH)₂ (109 mV dec⁻¹) and GNRs (287 mV dec⁻¹). EIS measurement was also used to determine the charge transfer resistance (R_{ct}) of the samples, with the GNRs/Co(OH)₂ showing the lowest resistance of 24 X, while the resistance of α -Co(OH)₂ alone was 150 X. The long-term durability of the GNRs/ Co(OH)₂ catalyst was also tested by the chronoamperometry method under a constant potential and found to be very stable for the OER process. In conclusion, the results of the study suggested that the GNRs/Co(OH)2 composite material is a highly active and stable catalyst for the OER process, with the synergetic effect between GNRs and Co(OH)2 leading to the enhancement of OER activity. It also highlights the importance of the morphology and state of Co(OH)₂ nanoparticles in the OER activity, with thin and well-dispersed nanoparticles on the GNRs leading to more exposed surface active sites and better contact with the substrate. Similar results have been reported by Pan et al. 138 that the a-Co(OH)2-NS electrode has a faster electron transfer process during the OER compared to α-Co(OH)₂-NF and bare NF, as indicated by the smaller charge transfer resistance (R_{ct}) value. The ECSA of α -Co(OH)₂-NS is also higher than those of α -Co(OH)₂-NF and bare NF. Additionally, α -Co(OH)₂-NS showed good long-term stability and negligible changes in current density at high potentials and high current densities.

Ge et al. 92 synthesized benzoate anion containing Co(OH)2 and demonstrated an overpotential of only 291 mV at a current density of 50 mA cm⁻² which is far better than that of Co(OH)₂ with CO₃²⁻ as interlayer anions. The enhanced activity was due to the increase in the interlayer spacing because of the large size of the benzoate anion as compared with that of the ${\rm CO_3}^{2-}$

anion. We concluded that the increase in the interlayer spacing added two attributes to the Co(OH)2; one is the availability of more exposed active sites and other is the increase in the ECSA evident from the increase in the $C_{\rm dl}$ of 1.43 mF cm⁻² for carbonate intercalated Co(OH)₂ to the 1.61 mF cm⁻² for benzoate intercalated Co(OH)2. Both of these factors have a significant contribution to the enhanced OER activity. The α -Co(OH)₂ nanosheets exhibit promising potential as oxygen evolution reaction (OER) electrocatalysts, holding considerable practical implications across various industries. Their unique structural properties, characterized by ultrathin, singlecrystalline features, suggest high efficiency in catalyzing OER processes. These nanosheets could find applications in energy storage systems, particularly in the development of more efficient and sustainable electrochemical devices such as watersplitting technologies. Additionally, their uniformity and welldefined structures pave the way for advancements in catalyst design, enhancing the performance and durability of electrocatalytic systems in industrial settings, driving forward innovations in renewable energy production and storage.

4.2.2 α-Co(OH)₂ in other morphologies as OER electrocatalysts. $\alpha\text{-Co(OH)}_2$ nanomeshes have been found to have superior electrocatalytic activity for the OER compared to bulk α -Co(OH)₂ and α -Co(OH)₂ nanosheets. In this study, the singular synthesis of ultrathin α-Co(OH)₂ nanomeshes is unveiled, marking the pioneering demonstration of this process. By subjecting cobalt salt to 2-methyl imidazole (MIM) at room temperature, a one-step reaction initiates the formation of these nanomeshes. A distinctive attribute of MIM surfaces is that it serves a dual purpose: functioning both as an alkali agent, yielding α-Co(OH)2, and as an etching reagent, essential in crafting mesopores within the α -Co(OH)₂ nanosheets. This dual role emerges through its coordination with cobalt, obviating the necessity for supplementary etching agents or procedures typically required in conventional post-treatment methodologies. α-Co(OH)2 nanomeshes have a low overpotential of 303 mV for a current density of 10 mA cm⁻², and a high mass activity of 31.3 A g⁻¹ when applying an overpotential of 303 mV. Additionally, the α-Co(OH)₂ nanomeshes have a smaller Tafel slope, indicating superior reaction kinetics, as well as smaller charge transfer resistance and larger electrochemically active surface area. The stability of the α-Co(OH)₂ nanomeshes is also good, as it can maintain 95% of its current density after 20 000 s of chronopotentiometric electrolysis testing. The improved performance of α-Co(OH)₂ nanomeshes is attributed to their abundant mesopores, which expose more active sites. XANES analysis of the α-Co(OH)₂ nanomeshes showed an increase in the Co valence state from Co²⁺ to Co³⁺ and the XAFS showed a decrease in intensity for the Co-O peak, indicating more oxygen vacancies in the exposed Co octahedrons.

An interesting study done by Zha et al. 140 discussed the use of hollow nano-dodecahedrons of α -Co(OH)₂ for OER activity. To synthesize α-Co(OH)2 and Cl-doped α-Co(OH)2 hollow nanododecahedrons, the procedure was initiated by obtaining ZIF-67 dodecahedrons in advance, following the methodology outlined in ref. 17 with specific modifications. Initially, a solution

comprising 1.43 g (~4.91 mmol) of Co(NO₃)₂·6H₂O and 3.24 g (~39.5 mmol) of 2-methylimidazole in 100 mL methanol underwent magnetic stirring at room temperature for 2 hours. The resultant solid product, obtained via centrifugation and subsequent methanol washes, was vacuum dried at 60 °C. Subsequent steps involved the addition of 20 mg ZIF-67 and 150 mg hexamethylenetetramine (HMT) into a 30 mL watermethanol mixture (with a volume ratio of 9:1) in the presence or absence of 50 mg NaCl. Following a 5 minutes stirring period, the system underwent thermal treatment at 40 °C for 4 hours within a Teflon-lined stainless-steel autoclave. After natural cooling to room temperature, a green precipitate was collected, washed thoroughly with deionized water and ethanol, and dried under vacuum conditions at 60 °C. The catalysts were activated by 10 CV cycles at 100 mV s⁻¹. The results showed that the Cldoped α-Co(OH)2/GCE had the most favorable OER kinetics and the highest electrocatalytic activity among the various catalysts tested, with a Tafel slope of 57 mV dec⁻¹ and a current density of 110.7 mA cm⁻². This is due to the transformation from Co(OH)₂ to CoOOH that occurs during the activation process. The hollow nano-dodecahedrons were synthesized through a hydrothermal method and were used as a substrate for the catalysts. The hollow structure of the nanododecahedrons is composed of a thin wall of the material and a large internal void. This structure provides a large surface area for the catalysts to adsorb and increases the number of active sites available for the electrocatalytic reactions, which in turn enhances the activity. Additionally, the hollow structure allows for a higher mass transport rate for the electrolyte which is important for the OER, since it involves the transfer of oxygen ions from the electrolyte to the electrode. This allows for a more efficient transfer of oxygen ions, leading to a higher current density. Furthermore, the hollow structure also provides a suitable environment for the catalysts to grow on, which can affect the stability and durability of the catalysts. The large internal void can also provide space for the formation of intermediate products and reduce the possibility of poisoning the catalysts, which can improve the stability and durability of the catalysts. Therefore, the use of the hollow nano-dodecahedrons as a substrate for the catalysts not only enhances the electrocatalytic activity but also improves the stability and durability of the catalysts which is important for long-term use. A similar study was performed by Liu et al.141 discussing the use of different types of Co(OH)2 as electrocatalysts for the OER. Different types of Co(OH)2 were synthesized by homogeneous precipitation of dilute CoCl2 solutions with HMT in the presence and absence of NaCl. α-Co(OH)2-Cl, which contains chloride anions, had superior OER performance compared to α- $Co(OH)_2$ -DS and β -Co(OH)₂, with a small onset overpotential of \sim 240 mV and a high current density of 10 mA cm⁻² at an overpotential of ~320 mV. α-Co(OH)₂-Cl had a more efficient kinetics for the OER and had the largest active surface area. The enhanced activity of the electrochemically etched α-Co(OH)₂-Cl is increased not just by the ECSA, but also by the intrinsic activity after electrochemical etching which plays a more important role. Overall, the study suggests that α -Co(OH)₂-Cl is a promising electrocatalyst for the OER in water splitting.

Bose et al. 142 described a study on the electrocatalytic OER of a catalyst, Y-CoII(H₂O)₆, by coating it with carbon on an electrode surface and conducting experiments in 0.1 M KOH (pH 13). The α -Co(OH)₂ film, which forms on the zeolite surface in the presence of KOH solution, plays a crucial role in boosting the OER. The structure of the electrode coating was observed using FESEM, which showed that the zeolite particles were well separated from each other, but the surface texture of the zeolite became corrugated with the appearance of α-Co(OH)₂ film/ sheet. The average size of the particles is approximately 1 mm. The α -Co(OH)₂ film plays a crucial role in promoting the OER by providing surface-bound redox-active species. Overall, the α -Co(OH)₂ film, which forms on the zeolite surface in the presence of KOH solution, plays a crucial role in boosting the OER by providing a smooth and uniform surface texture of the zeolite, promoting surface-bound redox-active species, and promoting higher-valence Co-species.

 $Co(OH)(C_6H_5COO) \cdot H_2O$ nanobelts are a new type of catalyst for the OER and have excellent performance compared to other Co-based catalysts, as well as benchmark catalysts such as RuO2.143 The nanobelts are formed by the incorporation of benzoate anions (C₆H₅COO⁻) into the Co(OH)₂ lattice, creating a unique catalyst structure. One key factor in the improved OER activity of these nanobelts is the large ECSA. The ECSA of the Co(OH)(C₆H₅COO)·H₂O nanobelts was calculated to be 19.5 μF cm⁻², which is significantly larger than that of the Co(OH)₂ (13.0 μF cm⁻²) and Co₃O₄ (14.0 μF cm⁻²) counterparts. This large surface area allows for more active sites for the OER to occur which is reflected in the higher OER current observed for the nanobelts. Another important factor is the relatively low overpotential required for the nanobelts to reach a current density of 10 mA cm⁻². The overpotential of the nanobelts was 360 mV, which is significantly lower than that of the Co(OH)₂ (440 mV) and Co₃O₄ (387 mV) counterparts. This means that less energy is required to drive the OER, making the nanobelts a more efficient catalyst. The Tafel slope was also found to be lower for the nanobelts (76 mV dec⁻¹) than for the Co(OH)₂ $(118 \text{ mV dec}^{-1})$, Co_3O_4 $(123 \text{ mV dec}^{-1})$ and Pt/C $(128 \text{ mV dec}^{-1})$ catalysts, but slightly higher than that of the benchmark catalyst RuO_2 (58 mV dec⁻¹). This indicates that the rate of the OER on the nanobelts is more favorable compared to that on other Cobased catalysts and Pt/C, but slightly slower than that on the benchmark RuO2. Overall, the incorporation of benzoate anions into the Co(OH)₂ lattice to form the Co(OH)(C₆H₅COO)·H₂O nanobelts results in a unique catalyst structure with large electrochemical surface area, low overpotential, and favorable kinetics for the OER, making them a promising catalyst for the OER. Similarly, Liu et al.144 described a new method for the synthesis of α-Co(OH)₂ nanoplates using Al³⁺ metal ions and ammonium solution, and the electrochemical performance of these nanoplates as electrode materials for supercapacitors and as OER electrocatalysts was measured. The OER of α-Co(OH)₂ nanoplates was found to be significantly better than that of β-Co(OH)2 particles, with a smaller onset potential and faster current increase. The mass activity and TOF of α-Co(OH)2 nanoplates were also found to be much higher than those of β-Co(OH)₂ particles. The mass activity of α-Co(OH)₂ nanoplates

was found to be 28.2 A g⁻¹ at a quite small overpotential of 348 mV, while it was only 0.2 A g^{-1} for β -Co(OH)₂ particles. The overpotential of 449 mV at 10 mA cm⁻² for β-Co(OH)₂ particles was evidently higher than that of α -Co(OH)₂ nanoplates. The α -Co(OH)₂ nanoplates also exhibit a linear increase in TOF with overpotential, affording much higher TOFs than β-Co(OH)₂ particles. EIS characterization was used to determine that the electronic properties of α -Co(OH)₂ nanoplates are related to the improved OER activity. The transport resistance of α-Co(OH)₂ nanoplates was found to be decreased.

The following conclusions could be drawn depending on the properties of intercalated anions in α -Co(OH)₂.

- Chlorides: chloride ions are known to increase the conductivity of the electrolyte and lower the overpotential for the OER. This is because chloride ions increase the electron transfer rate by facilitating the formation of a stable and conductive film on the surface of the electrode. However, at high concentrations, chloride ions can also lead to corrosion and dissolution of the Co(OH)2, which can negatively affect the performance of the OER.
- Carbonates: carbonate ions are known to improve the OER performance by providing a basic environment that stabilizes the Co(OH)2 and increases the pH of the electrolyte. This can increase the activity of the Co(OH)2 and lower the overpotential for the OER. However, at high concentrations, carbonates can also lead to carbonate-induced dissolution of the Co(OH)₂, which can negatively affect the performance of the OER.
- Nitrates: nitrate ions are known to increase the conductivity of the electrolyte and lower the overpotential for the OER. This is because nitrate ions increase the electron transfer rate by facilitating the formation of a stable and conductive film on the surface of the electrode. However, at high concentrations, nitrate ions can also lead to corrosion and dissolution of the Co(OH)2, which can negatively affect the performance of the OER.
- Phosphates: phosphate ions are known to improve the OER performance by increasing the stability of the cobalt hydroxide in aqueous environments and providing a basic environment that stabilizes the Co(OH)2 and increases the pH of the electrolyte. This can increase the activity of the Co(OH)2 and lower the overpotential for the OER.
- Benzoates: benzoate ions are known to improve the OER performance by increasing the stability of the cobalt hydroxide in aqueous environments and providing a basic environment that stabilizes the Co(OH)2 and increases the pH of the electrolyte. This can increase the activity of the Co(OH)₂ and lower the overpotential for the OER.

It is important to note that the specific effects of these interlayer anions on the OER in α -Co(OH)₂ will depend on the concentration of the anions and the specific conditions of the reaction. Additionally, the effect of interlayer anions on the OER can be different depending on the type of Co(OH)2, the surface area, particle size and morphology of the Co(OH)₂, and the potential and temperature of the reaction. Therefore, it is important to conduct a detailed study of the OER in the presence of these interlayer anions to understand their specific effects and optimize the performance of the reaction. It is also

important to note that this information is based on the current research available, and new discoveries and developments may occur in the future which may change the current understanding of the specific properties and trends.

4.3 Cobalt oxyhydroxide (CoOOH)

In the subsequent sections we will discuss the impact of different morphologies of α -Co(OH)₂ on the oxygen evolution reactions.

4.3.1 Cobalt oxyhydroxide (CoOOH) as an OER electrocatalyst. The electrocatalytic activity of CoO_xH_v/CC is predominantly attributed to CoO_xH_v NSAs. 145 It was found that the CoO_xH_{\(\nu\)}/CC only requires an overpotential of 430 mV to deliver a current density of 10 mA cm⁻², which is far superior to that of RuO₂/CC and CoO_xH_y-NO/CC (630 and 820 mV, respectively). The Tafel slope for CoO_xH_v/CC (121 mV dec⁻¹) is also far lower than those of CoO_xH_v -NO/CC (377 mV dec^{-1}) and RuO_2 /CC (394 mV dec⁻¹), indicating much faster catalytic kinetics enabled by the CoO_xH_v/CC electrode. These results demonstrate that the CoO_xH_y/CC outperforms RuO₂/CC, the current noble metal-based benchmark electrocatalyst for the OER, and thus requires much less energy for the electrochemical processes. The exceptional electrocatalytic OER activities of CoO_xH_v/CC electrodes are attributed to their unique nanostructures. The ECSA of the CoO_xH_v/CC electrode is estimated to be 154 mF cm⁻², far higher than those of RuO₂/CC (6.5 mF cm⁻²) and CoO₂H_v-NO/CC (25.2 mF cm⁻²) (Fig. 9). This high ECSA is a consequence of the ultrasmall grain size and excellent porosity of the CoO_xH_v NSAs, and thus can contribute significantly to the excellent electrocatalytic OER activities by facilitating the efficient exposure of abundant active sites for the OER.

Chen et al.146 showed that the Co/PANI HNSs are a hierarchical porous hybrid material composed of cross-linked PANI and ultrathin CoOOH nanosheets. They were fabricated by combining a wetness-impregnation method and a subsequent electrochemical process. LSV was performed to evaluate the OER activities of PANI, COOOH, and Co/PANI HNSs. The results showed that the Co/PANI HNS catalyst has the lowest onset potential and highest anodic current density among the three samples, even better than those of the reference catalyst of IrO₂. The Co/PANI HNSs only require 291 mV overpotential to reach a current density of 10 mA cm⁻², significantly lower than those of PANI (474 mV), CoOOH (407 mV), and even IrO2 (344 mV). The catalytic performance of Co/PANI HNSs is comparable or even superior to that of some recently reported state-of-the-art OER electrocatalysts based on CoOOH-type active species under alkaline conditions. The exceptional performance is attributed to the hierarchical porous hybrid material composed of cross-linked PANI and ultrathin CoOOH nanosheets, which is conducive to the adsorption, activation, and conversion of the reactants and also the large ECSA of the material. The CoOOH nanosheets are strongly bound to PANI through the formation of Co-N coordination bonds, which act as a structural bridge between the two components. This not only improves the stability of the structure but also reduces the resistance to

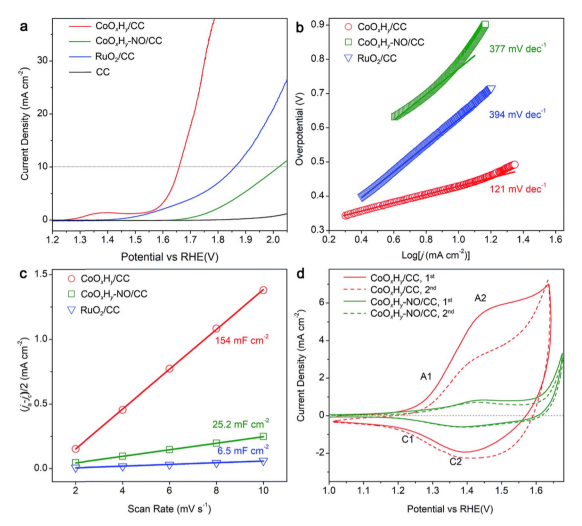


Fig. 9 (a) Polarization curves of COO_xH_v/CC , COO_xH_v-NO/CC , RuO_2/CC and CC in 0.1 M PBS (iR-corrected, 2 mV s⁻¹). (b) The corresponding Tafel plots. (c) Capacitive current densities at 1.25 V versus RHE as a function of scan rate for COO_xH_v/CC , COO_xH_v-NO/CC and RuO_2/CC . (d) CVs of CoO_xH_v/CC and CoO_xH_v-NO/CC in 0.1 M PBS (iR-corrected, 5 mV s⁻¹). This figure has been reproduced from ref. 145 with permission from Royal Society of Chemistry, copyright 2019.

charge transfer at the interface. The interaction between Co dorbitals and PANI's π-conjugated ligands in Co/PANI HNSs allows for efficient electron transfer from CoOOH to PANI, which improves the adsorption energy of reactants and products during the OER process. Additionally, the presence of mesopores and macropores in Co/PANI HNSs allows for easy access of electrolytes to active sites, leading to faster reaction kinetics. These factors contribute to the exceptional performance of Co/PANI HNSs as an OER electrocatalyst.

Meng et al. 147 fabricated laser induced oxygen vacant CoOOH for performing the OER. LSV curves were obtained at a scan rate of 5 mV s⁻¹, and bulk CoOOH (B-CoOOH) and commercial RuO₂ were used as references. The results showed that L-CoOOH required a lower overpotential of 330 mV to reach a current density of 10 mA cm⁻², which is better than those of RuO₂ (340 mV) and BCoOOH (410 mV). Additionally, the Tafel slope of L-CoOOH (63.2 mV dec⁻¹) is lower than that of B-CoOOH (75.4 mV dec⁻¹) and close to that of RuO₂ (64.9 mV dec⁻¹), indicating favorable OER kinetics for L-CoOOH. After 10

hours of testing, RuO₂ only maintained 69% of its initial current density, while L-CoOOH retained 95% of its initial current density. The morphology and composition of L-CoOOH remained unchanged after the OER durability test. The $C_{\rm dl}$ value of L-CoOOH was found to be 7.4 times higher than that of B-CoOOH, indicating the existence of more active sites in L-CoOOH. The electrical conductivity of L-CoOOH was also higher than that of B-CoOOH due to a smaller R_{ct} . Finally, the study used X-ray photoelectron spectroscopy (XPS) to explore the surface valence state of L-CoOOH and B-CoOOH. The XPS results showed that L-CoOOH had a higher ratio of Co²⁺ to Co³⁺ than B-CoOOH, which was attributed to the increased active sites and faster electron transfer in L-CoOOH. Overall, it is concluded that L-CoOOH is a promising OER catalyst with excellent performance and durability and that its superior performance is due to the increased active sites and faster electron transfer.

The research conducted by Wang et al.148 described a new method for synthesizing hierarchical ultrathin sheet-based

CoOOH hollow nanospheres using a room-temperature selftemplate strategy. The researchers used Pi-doped Co-glycerate hollow nanospheres as the self-sacrificing templates, which were then converted to CoOOH hollow nanospheres through an outside-in pathway involving ionization, anion-exchange reaction, and oxidation in an alkaline environment. To maintain the spherical structure, the precursors were coated with a Nafion membrane, which was easily removed by washing with ethanol. The resulting CoOOH hierarchical hollow nanospheres have several desirable properties, including ultrathin nanosheets and abundant oxygen vacancies, which provide abundant catalytic active sites, shorten charge/ion diffusion paths, reduce charge-transport resistance, and enhance the stability for the OER. Therefore, the CoOOH hierarchical hollow nanospheres obtained by this method have extraordinary electrocatalytic activity and good stability for the OER. The hierarchical CoOOH hollow nanospheres have been shown to have high catalytic activity in the OER, with an overpotential as low as 275 mV at a current density of 10 mA cm $^{-2}$. It is superior to the γ -CoOOH nanosheets (overpotential of 304 mV) and the bare Ni foam (overpotential of 362 mV) at the same current density. The hierarchical CoOOH hollow nanospheres also have lower charge-transfer resistance and faster charge transfer kinetics than the γ-CoOOH nanosheets. Additionally, the hierarchical CoOOH hollow nanospheres have a higher electrochemical double-layer capacitance (22.6 mF cm⁻²) than the γ-CoOOH nanosheets (4.6 mF cm⁻²), indicating more active sites on the surface of the hierarchical CoOOH hollow nanospheres. The hierarchical CoOOH hollow nanospheres also showed longterm stability, with no change in polarization curves even after 1000 cycles. However, the crystalline structure of the hierarchical CoOOH hollow nanospheres was not preserved after the OER test.

Ye et al.149 investigated the electrocatalytic activity of a material termed Fe_{0.33}Co_{0.67}OOH PNSAs/CFC as an oxygen evolution reaction (OER) catalyst. The material exhibited the highest OER electrocatalytic activity among the materials tested, which included commercial IrO2 coated on Ni foam (IrO2/NF) and CoOOH PNSAs/CFC. The Fe/Co molar ratio in the Fe_{0,33}Co_{0,67}OOH PNSAs/CFC material was controlled by adjusting the time of in situ anodic oxidation of α -Co(OH)₂ in a solution of 0.01 M $(NH_4)_2$ Fe $(SO_4)_2$. The study found that the Fe/Co molar ratio in the material can be increased to about 1:2 and remains constant after 30 minutes, and the catalysts composed of Fe/Co = 0.493 (Fe_{0.33}Co_{0.67}OOH PNSAs/CFC) exhibit the highest OER electrocatalytic activity. The Fe_{0.33}-Co_{0.67}OOH PNSAs/CFC material required a much lower overpotential than the other materials tested, specifically, 266 mV at 10 mA cm⁻², whereas CoOOH PNSAs/CFC and IrO₂/NF catalysts required overpotentials of about 331 and 290 mV, respectively. The Fe_{0.33}Co_{0.67}OOH PNSAs/CFC also showed much higher mass electrocatalytic activity for the OER than CoOOH PNSAs/ CFC and IrO₂/NF. The Tafel slope of Fe_{0.33}Co_{0.67}OOH is only about 30 mV dec⁻¹, which is much lower than those of CoOOH PNSAs/CFC (56.4 mV dec^{-1}) and IrO_2/NF (77.2 mV dec^{-1}). The EIS of Fe_{0.33}Co_{0.67}OOH PNSAs/CFC and CoOOH PNSAs/CFC, and the corresponding circuit model fitting analysis were also

performed. The EIS results showed that the R_{ct} of Fe_{0.33}Co_{0.67}-OOH PNSAs/CFC was smaller than that of CoOOH PNSAs/CFC, suggesting a fast electron transfer is realized in Fe_{0.33}Co_{0.67}OOH PNSAs/CFC, which would expedite the OER and reduce the overpotential due to its high conductivity. The study also measured the TOF of the electrocatalysts, which showed that the TOF of Fe_{0.33}Co_{0.67}OOH PNSAs/CFC was much higher than that of CoOOH PNSAs/CFC, further illustrating the improved activity of the material.

Bergmann et al.150 used DFT calculations to explain the change in site occupancy of Co²⁺ Td ions to empty Oh sites with the μ-O vacancy formation in the reaction zone during the OER, as calculated for the Co₃O₄(100) surface. Oxygen vacancies also probably migrate into the subsurface; thus, the authors consider this process to induce the restructuring of the nearsurface at elevated O2 evolution rates. Oxygen vacancy formation on the $Co_3O_4(110)$ surface has less impact on the local atomic structure. XANES was used to determine the change in the cobalt K XANES spectra, which suggests oxidation of the metal ion to Co4+. The lower charge transfer energy in $\text{Co}^{3+}/^{4+}\text{O}_x(\text{OH})_v$ also allows the formation of hole density on the oxygen ligand, as identified for IrO_x(OH)_y. Tafel slopes decreased as the mean electrochemical reducibility of the Co3+-O sites increased, providing evidence that this reducibility is closely correlated with favorable electrocatalytic properties. Better electrocatalytic properties with a more pronounced Co²⁺/ Co³⁺ redox transition was also identified under phosphate-free, alkaline reaction conditions. Additionally, the redox charge of Co²⁺/Co³⁺ is a better descriptor of the OER active site density than the overall number of redox sites. In summary, the study found that the structural transformation of Co2+ oxides into Co^{3+} Oh-containing $CoO_x(OH)_y$ during OER conditioning occurred in the near-surface region, leading to a structurally unified catalytically active state with a higher mean cobalt oxidation state and a contracted CoO₆ bonding environment. DFT calculations showed that this transformation was caused by the formation of oxygen vacancies in the reaction zone. The electrocatalytic properties of CoO_x(OH)_v were found to be correlated with the near-surface electronic structure and degree of defects. The Tafel slopes were found to decrease as the mean electrochemical reducibility of the Co3+-O sites increased, indicating that this reducibility is closely correlated with favorable electrocatalytic properties.

Cui et al.151 studied the promotion of the OER via the synergistic effect of coupling CoOOH with MnOOH. The MnOOH is a well-known OER catalyst because of its high activity and stability. It has been shown to have a low overpotential and small Tafel slope, indicating fast OER kinetics. However, the activity of MnOOH can be further improved by synergistically coupling it with other catalysts, such as cobalt oxide (CoOOH). B-MCO^{-0.1} catalyst was found to have a lower overpotential ($\eta 10 = 313 \text{ mV}$) and smaller Tafel slope (87 mV dec⁻¹) compared to control samples such as γ-MnOOH (494 mV and 122 mV dec^{-1}) and CoOOH (420 mV and 140 mV dec^{-1}). This suggests that the interfacial interaction between the γ -MnOOH and CoOOH particles in the B-MCO^{-0.1} catalyst results in faster OER kinetics. The study also found that the B-MCO^{-0.1}

had the highest mass activity (4.3 A mg^{-1}) and TOF (50.3 s^{-1}) among the control samples. These results indicate that the interfacial interaction between γ-MnOOH and CoOOH in the B-MCO^{-0.1} catalyst not only improves the activity, but also the stability of the surface species. One possible mechanism for the synergy between MnOOH and CoOOH is that the CoOOH particles act as a conductive matrix, facilitating electron transfer during the OER process. The γ-MnOOH particles, on the other hand, provide a large number of active sites for the OER to occur. The strong interfacial interaction between these two catalysts results in a higher number of active sites per unit area and faster electron transfer, leading to improved OER activity. Additionally, as seen in the study, the B-MCO^{-0.1} also showed the biggest $C_{\rm dl}$ value (11.5 mF cm⁻²) and the largest ECSA among the control samples, indicating more accessible active sites in the OER; both of these factors also contribute to the improved activity of the B-MCO^{-0.1} catalyst.

In a similar study, Xu et al. 152 evaluated the OER electrochemical performance of a NiSe@CoOOH core-shell structure as a 3D integrated electrode in a 1 M KOH aqueous electrolyte. The NiSe@CoOOH core-shell structure was found to have superior OER activity compared to bare Ni foam, NiSe NWs/NF, and RuO2/NF electrodes. The NiSe@CoOOH core-shell structure had a sharp onset potential at 217 mV with greatly enhanced OER current, requiring an overpotential of only 292 mV to reach 100 mA cm⁻². The Tafel slope of the NiSe@-CoOOH core-shell structure electrode was approximately 65 mV dec⁻¹, which is lower than that of the other electrodes tested, indicating that the core-shell structure accelerates the catalytic kinetics for the OER. The authors suggest that the as-prepared NiSe@CoOOH/NF provides the real catalytic mediator CoOOH nanosheets with more surface-active sites for the OER and the core-shell structure accelerates the charge transfer during the OER. The unique structure of the NiSe core and the coating with 200 nm CoOOH nanosheets may improve the structural stability and accelerate fast electron transfer which is beneficial for fast transportation of electrons from the NiSe core to the CoOOH shell. Additionally, the stability of the electrocatalyst was found to be good, showing consistent results even after 12 hours of electrolysis.

Moreover, Liu et al. 153 put in some effort to unveil the role of Co³⁺ in the OER with CoOOH based materials. The OER was evaluated in an O_2 -saturated 1 M KOH (pH \sim 14). The results of LSV measurements with a sweep rate of 5 mV s⁻¹ showed that the Co³⁺-33 h sample exhibited prominently enhanced catalytic activity for the OER compared to other Co³⁺ samples and β- $Co(OH)_2$. The low overpotential of 395 mV at 10 mA cm⁻² indicated its good catalytic activity towards the OER and is superior to many recently developed catalysts. The enhanced activity was attributed to the unique Co³⁺-rich surface composition (Co(OH)₂/CoOOH) of Co³⁺-33 h, in which the Co³⁺ ions served as the active sites for the OER. The varied charge distribution, as evidenced by the negative shift of Co $2p_{3/2}$ in Co^{3+} 33 h compared with Co(OH)2, might result in a lower energy barrier for the adsorption of H2O during water oxidation. Furthermore, the varied catalytic activity between Co³⁺-24 h, Co³⁺-33 h, and Co³⁺-42 h can be related to the content of Co³⁺ on

the surface, as revealed by the XPS analysis. The constant current activation time increased, more active Co³⁺ ions were generated and the content of Co3+ reached a maximum after activation of 33 h. In summary, the enhanced OER activity of the Co³⁺-33 h sample was due to the unique Co³⁺-rich surface composition (Co(OH)₂/CoOOH) of Co³⁺-33 h, in which the Co³⁺ ions served as the active sites for the OER, and the varied charge distribution, which resulted in a lower energy barrier for the adsorption of H2O during the OER, leading to improved OER performance.

Reikowski et al. 154 discussed the preparation of pure phase epitaxial Co₃O₄(111) and CoOOH(001) thin layers on Au(111) through electrodeposition. The stability of the CoOOH(001) layer and the restructuring of the Co₃O₄(111) spinel oxide near the surface region are discussed, as well as the catalytic properties of both layers. The stability of the different cobalt oxides is defined by the following redox reactions:

$$3\text{Co(OH)}_2 + 2\text{OH}^- \rightarrow \text{Co}_3\text{O}_4 + 4\text{H}_2\text{O} + 2\text{e}^-$$

$$\text{Co}_3\text{O}_4 + \text{OH}^- + \text{H}_2\text{O} \rightarrow 3\text{CoOOH} + \text{e}^-$$

$$\text{CoOOH} + \text{OH}^- \rightarrow \text{CoO}_2 + \text{H}_2\text{O} + \text{e}^-$$

Conclusively, this study highlighted the stability and structural changes of the CoOOH(001) thin layer on Au(111) when subjected to different potentials. According to thermodynamic, the CoOOH film exhibits a wide stability range and does not exhibit structural changes down to 0.4 V. The stability of the CoOOH layer is attributed to its smooth surface, which is composed of close-packed (001)-terminated terraces. The absence of structural changes in the OER range suggested that the CoOOH lattice remains unchanged in this range and the natural (001) cleavage plane is stable and active during the OER. DFT calculations support this observation, showing that the CoOOH(001) surface termination gradually changes from OH to O in this potential regime, and reaches a fully O-terminated surface at 1.5 V vs. RHE. Additionally, DFT calculations also exclude the formation of bulk crystalline CoO2, even at high potentials where its formation should be thermodynamically favorable. These findings provide a deeper understanding of the stability and activity of the CoOOH(001) thin layer as a potential electrocatalyst in OER applications. Potential-dependent restructuring of Co₃O₄ film surfaces refers to the changes in the surface structure of a Co3O4 film that occur at different applied potentials. The article discussed the formation of a "skin layer" on the surface of the Co₃O₄ film that occurs at potentials between 1.2 and 1.67 V. This skin layer is a reversible, ≈ 1 nm thick layer that forms on the top and side walls of the Co₃O₄(111) grains. The formation of this skin layer is not directly related to the catalytic reaction (OER) as previously assumed, but rather reflects the thermodynamics of the Co₃O₄/ CoOOH phase transition. It is proposed that the formation of this skin layer is caused by the displacement of Co²⁺ cations from tetrahedral to octahedral symmetry, as the Co oxidation state increases from Co²⁺/Co³⁺ to Co³⁺ for potentials above

Proposed OER mechanism. This figure has been reproduced from ref. 155 with permission from American Chemical Society, copyright Fig. 10 2020.

1.2 V, which is well below the thermodynamic limit of the OER required to initiate the reaction. Absolutely, the formation of this "reversible skin layer" seems to be linked more to the chemical changes within the oxide itself rather than being directly tied to the OER. This means that the transformation of the material from, say, Co₃O₄ to CoOOH and back, which forms this skin layer, is happening independently or prior to the onset of the OER process.

The charge transfer that occurs during this specific chemical transformation contributes significantly to what's known as pseudocapacitive current. This current, observed before the OER, stems from this transformation process within the material rather than the electrochemical reactions involved in oxygen evolution. Understanding and harnessing this reversible skin layer formation could potentially lead to advancements in pseudocapacitive-based energy storage or electrochemical devices. This restructuring requires significant transport of oxygen and Co ions through the oxide lattice, which can occur rapidly due to the low thickness of the skin layer. The skin layer does not form on a perfectly smooth Co₃O₄(111) surface but rather on nanoscale 3D islands that expose other facets to the electrolyte, which may facilitate the layer formation by providing channels for cation migration within the oxygen framework. The catalytic properties of epitaxial samples of Co₃O₄(111) and CoOOH(001) films indicate that a smooth epitaxial CoOOH(001) layer is nearly as good a catalyst as a Co₃O₄(111) layer, which is in contrast with general

assumptions in the literature. The study also suggests that the OER active sites in the films are di-μ-oxo-bridged ion clusters in amorphous CoCat catalyst films and μ3-O/OH sites on the (001) plane of CoOOH. The findings of the study may have significant implications for the understanding of the nature of OER active sites on Co oxide catalysts and suggest that well-defined epitaxial films may be useful in future studies. In conclusion, the electrochemical charge (Q0) measured in the pre-OER region for CoOOH(001) is attributed to adsorption/desorption on the oxide surface, specifically protonation/deprotonation of the outermost oxygen atoms on the top surface plane and within the grain boundaries. In the case of Co₃O₄, the charge Q0 corresponds to a charge transfer that involves the bulk of the Co₃O₄ epitaxial film. This is supported by optical reflectivity measurements and previous observations for CoCat. The changes in the bulk charge state are linked to reversible changes in unit cell volume. However, the nature of the charges associated with the substantial pseudocapacitive charge/discharge in Co₃O₄ is still not clear, and more work is necessary to check the hypothesis of intercalation of solution species. It is important to note that the current in the pre-OER range cannot be easily employed to estimate the electrochemical surface area or redoxactive surface sites, as dividing the pseudocapacitance of the oxide by a standard double layer capacitance leads to an erroneous ECSA that is several orders of magnitude larger than derived from independent structural characterization.

The mechanism proposed by the group of Xile Hu155 in Fig. 10 for the OER on Co-based catalysts is consistent with spectroscopic and electrokinetic data. The mechanism suggests that the oxidation of Co(III) to Co(IV) is reconciled by considering a di-Co active site, where the oxidation of the first Co(III) ion is accompanied by a loss of 2 protons, and the oxidation of the second Co(III) ion is accompanied by a loss of 1 proton. This results in the formation of a Co(IV)-O' intermediate, which is proposed to be in equilibrium with a bridging Co-superoxide-Co species. The mechanism proposed in Fig. 10 predicts that the concentration of the superoxide species would increase with increasing potentials, which is supported by the observed data. The potential-determining step is proposed to be the oxidation of the second Co(III) to Co(IV) ion. It is different from previous mechanisms proposed for Co-based catalysts, which mostly invoke Co(IV)=O or Co(III)-O' as the highest valence Co species responsible for O-O bond formation.

- The first step in the proposed mechanism is the oxidation of the first Co(III) ion, which is accompanied by a loss of 2 protons. This results in the formation of a Co(III)- $(\mu$ - $O)_2$ -Co(IV)species. The oxidation of the second Co(III) ion is accompanied by a loss of 1 proton and results in the formation of a Co(III)-(μ-O)2Co(IV)-OH species. These two steps represent the protoncoupled electron transfer (PCET) steps in the mechanism.
- The next step in the mechanism is the oxidation of the resting state CoO₂ by 1 electron. This occurs at the O anion to give a Co(iv)-O' species, which is proposed to be in equilibrium with a bridging Co-superoxide-Co species. This species is the active oxygen species observed in the Raman spectra. The oxidation of the resting state CoO₂ to the Co(IV)-O' species is considered a pre-equilibrium.

The rate-determining step (RDS) in the mechanism is the release of O2 from the Co-superoxide-Co species with a concomitant H2O binding. This step is followed by the binding of another H₂O and further 1-electron oxidation, which regenerates the starting Co(III) form of the catalyst (CoOOH). This sequence would give a Tafel slope, a first order in [OH⁻], as well as D-EIE and -KIE that agree with experimental values. The proposed mechanism is consistent with previous experimental and theoretical studies, which show that the formation of Co(IV)-O' is necessary for O-O bond formation catalyzed by various small Co-oxide fragments. A theoretical study suggested that assuming 4 consecutive PCETs as the mechanism of OER, the most active site of crystalline β -CoOOH has a resting state of Co(III) instead of Co(IV). However, such a mechanism would not apply to this system as the electrokinetic data cannot be fit by 4 PCETs. It is worth noting that dioxygen release as a ratedetermining step in the OER has not been proposed for Cobased catalysts, but it does have several precedents in other types of catalysts.

4.4 Doped-cobalt hydroxides as OER electrocatalysts

In recent years, researchers have been exploring different doping strategies to optimize the performance of doped cobalt hydroxide catalysts for the OER. For example, alloying cobalt hydroxide with other metals has been shown to improve the

catalytic activity and stability, and using different synthetic methodologies can result in different morphologies which can affect the catalytic properties. Furthermore, doped cobalt hydroxide can be used under both acidic and basic conditions, which makes it versatile for different types of applications. Also, it has a good performance at both low and high current density, this feature makes it suitable for different types of devices.

The study performed by Zhang et al. 156 discusses the evaluation of the catalytic performance of CCHH-based electrocatalysts for the oxygen evolution reaction (OER) in 1 M KOH using a three-electrode system. One of the main findings of the study is that the catalytic activities were greatly boosted after Fe doping. The optimal Fe-CCHH/NF-30 exhibited the best OER performance, requiring a much smaller overpotential of 200 mV to deliver a current density of 10 mA cm⁻². This means that it requires less energy to drive the reaction to the same current density compared to other catalysts, making it more efficient. The OER activity of Fe-CCHH/NF-30 was also found to be better than that of the commercial RuO2 with the same loading, and this is quite competitive when compared to other CCHH-based or Co-based catalysts. The study suggests that the superior OER performance of Fe-CCHH/NF-30 is due to the tailored electronic structure that favors the reaction kinetics. The XPS spectrum of Co 2p demonstrates a noticeable negative shift upon doping with Fe, indicating a lower oxidation state of Co compared to its undoped counterpart. This phenomenon contributes significantly to the in situ conversion of Co to higher oxidation states during electrochemical investigations, thereby augmenting its overall activity. The tailored electronic structure is likely a result of the atomically doped Fe, which can greatly enhance the intrinsic activity of CCHH. It is an important finding, as it suggests that single atom catalysts may be more effective than traditional catalysts that rely on multiple atoms to drive the reaction. The study also suggests that the approach of in situ electrochemical transformation may be a promising strategy to design single atom catalysts with high activity and stability. In situ electrochemical transformation involves introducing the dopant atoms (in this case Fe) during the electrocatalyst synthesis process, rather than adding them afterward. This allows for better control over the electronic structure of the catalyst, which in turn leads to better performance. Finally, the Fe-CCHH/NF-30 had excellent durability under long-time stability and large current density conditions. This is an important factor for practical applications, as catalysts that degrade quickly or lose activity over time would not be useful in real-world settings. However, further research is required to understand the potential of single atom catalysts in real-world applications and how to design these catalysts for better performance and stability.

Jin et al.157 suggested that the incorporation of Fe into Co(OH)₂ plays a crucial role in promoting the OER. The catalytic ability of Fe-incorporating $Co(OH)_2$ (α - $Co_4Fe(OH)_x$) is notably higher than that of pure $Co(OH)_2$ (α - $Co(OH)_2$). This indicated that Fe promotes the OER by increasing the catalytic activity of $Co(OH)_2$. Additionally, the polarization curve of α - $Co_4Fe(OH)_x$ revealed a small onset potential of 0.26 V for the OER, indicating that the Fe-incorporating Co(OH)2 catalyst is able to

initiate the OER at a lower potential compared to pure Co(OH)₂. This suggests that Fe may be facilitating the OER by making it easier for electrons to transfer to the catalytic sites, leading to a lower activation energy for the reaction. Furthermore, the α - $Co_4Fe(OH)_r$ catalyst also achieved a high TOF of 0.027 s⁻¹ at an overpotential of 0.3 V, which is 11 and 5 times higher than those of α -Co(OH)₂ and IrO₂, respectively. This suggests that the Fe incorporation increases the rate of OER by increasing the number of active sites available for the reaction to occur. In summary, the study suggested that Fe plays a significant role in promoting the OER by increasing the catalytic activity, decreasing the onset potential, and increasing the turnover frequency of the Co(OH)2 catalyst.

The LSV curves of the $Co(OH)_2$ and $Ni_xCo_y(OH)_2$ materials were measured at a scan rate of 10 mV s⁻¹ by Wang et al. 158 The enlarged LSV curve demonstrates the oxidation peaks of CoII/ CoIII (~1.15 V vs. RHE) and CoIII/CoIV (~1.43 V vs. RHE). When doping the Ni in this system (15%), the OER activity increases (overpotential = 383 mV). The $Ni_{0.25}Co_{0.75}(OH)_2$ exhibits the best OER performance with an overpotential of 352 mV. This is because the Ni_{0.25}Co_{0.75}(OH)₂ has the highest intrinsic activity for the OER, which is likely due to the specific electronic structures that are beneficial for the OH⁻ adsorption. The Ni doping leads to a positive shift of the oxidation peak of Co species, and the Ni_{0.25}Co_{0.75}(OH)₂ has the largest CoII/CoIII oxidation peaks compared to other samples. This suggests that Ni doping causes a change in the electronic structure of Co atoms, leading to the difference in OER behavior. Furthermore, the voltammetry of $Ni_xCo_v(OH)_2$ shows a strong dependence of the CoII/III potential on the Ni content, suggesting a strong electronic coupling between Ni and Co in the solid. The Ni_{0.25}Co_{0.75}(OH)₂ has the largest number of tri- and tetra-valent metal cations on the surface of the nanosheet material before the occurrence of the OER. Therefore, the Ni_{0.25}Co_{0.75}(OH)₂ has the smallest overpotential since these high-valence Co cations are regarded as active sites of the OER. This indicates that Ni_{0.25}Co_{0.75}(OH)₂ has a much larger active surface area compared to other samples. In summary, the doping of Ni into Co-based hydroxide materials enhances the OER activity by changing the electronic structures of Co atoms, leading to a positive shift of the oxidation peak of Co species and a stronger electronic coupling between Ni and Co. Similar results on the role of Ni in boosting the OER of Co(OH)2 could be found in the study by Guo et al. 159

In a similar study Zeng et al. 160 suggested that the addition of iron and nickel to cobalt hydroxide (Co(OH)2) improved the performance of the material as an electrocatalyst for the OER. The LSV data from the study suggests that the Co-Ni-Fe511 compound nanosheets have a higher polarization current and lower overpotential compared to Co(OH)2 and also show better performance than Co-Ni511 and Co-Fe511. The Tafel plots also indicate that the incorporation of Fe and Ni results in a more rapid OER rate and a smaller Tafel slope. Additionally, the electrochemical impedance spectroscopy results suggest that the Co-Ni-Fe511 catalyst has faster electrode kinetics and higher faradaic efficiency. Overall, it is suggested that the

synergistic effect of Fe and Ni dopants enhances the OER activity of cobalt hydroxide.

Cao et al. 161 synthesized α-Co(OH)2 using different solvents and also doped it with different transition metal ions (Ni²⁺, Cu²⁺, and Fe²⁺) to study their effect on the electrochemical performance of the material as an OER electrocatalyst. They used CV and LSV to measure the OER performance of the different samples. The results show that α -Co(OH)₂ synthesized in pure water exhibited poor OER performance, while the addition of Cu²⁺ or Fe²⁺ significantly reduced the overpotential, with the best performance being obtained when the samples contained 30% Cu²⁺ or 20% Fe²⁺ in a mole. In contrast, doping with Ni2+ did not improve the performance. However, it also contrasts with the initial studies where doping with Ni enhances the OER activity. They also found that Fe-Co(OH), not only had the lowest overpotential but also had excellent stability and the highest TOF value which was 3 times higher than that of α -Co(OH)₂. Additionally, Fe-Co(OH)₂ had the lowest Tafel slope, indicating its quicker electrocatalytic kinetics and higher OER electrochemical catalytic activity.162 Furthermore, EIS was conducted to evaluate the electron transport ability and the results show that Fe-Co(OH)2 and Cu-Co(OH)2 had smaller Rct values, indicating more efficient charge transport during the electrochemical OER process than α-Co(OH)2. A similar study on the role of Cu doping to increase the OER activity of Co(OH)2 is reported by Chen et al. 163

In an interesting study, electrochemical tests of pure and modified CoOOH nanowires were performed by Chen et al. 164 The results showed that the maximum current density of NiCoOOH-A (99% greater than pure CoOOH) and the potential required to attain 10 mA cm⁻² (decreased by 20 mV) were improved when compared to those of pure CoOOH. When the surface Ni concentration was increased, the OER activity also increased. Specifically, at a concentration of 9.7% Ni, the OER activity increased to 400% at 1.55 V when compared to pure CoOOH. However, when the surface Ni concentration was increased to 15%, the OER activity decreased from the optimal value. It was also observed that Ni-CoOOH-C with 15% surface Ni has lower activity than 5.5% Ni but exhibits higher activity than pure CoOOH. A comparison of the OER activity of Ni-CoOOH samples to NiO_xH_v nanoparticles illustrates that Nimodification of CoOOH creates more active OER sites than does CoOOH or NiO_xH_v alone. On the other hand, Mn modification showed no improvement or decrease in the OER activity of pure CoOOH at all levels of Mn incorporation. The apparent lack of influence of Mn incorporation on the OER activity can be explained by the formation of both inactive Mn4+ and active Mn³⁺ ions from Mn incorporation. It was recently suggested that accidental Fe doping may have been the reason for the improved OER activity of certain Co/Ni oxide or oxyhydroxide catalysts. However, the authors of this study believe that the trends in OER activity reported here are due to Ni and Mn incorporation as opposed to accidental Fe doping. Inspection of the Tafel regions in LSV of Ni- and Mn-CoOOH samples revealed a decrease in the Tafel slope for Ni-CoOOH compared to pure CoOOH, specifically from 53 mV dec⁻¹ to 36 mV dec⁻¹. However, no change was observed upon Mn modification.

Nevertheless, a contradictory study on Mn doping in CoOOH for boosting the OER was done by Huang et al. 165 The research presents the water oxidation performance of electrocatalysts made of Mn-confined CoOOH nanosheets. The electrocatalysts were evaluated using steady-state electrochemical measurements in 1 M KOH solution, specifically cyclic voltammetry (CV) and polarization curves. Mn-doped CoOOH nanosheets showed improved OER performance compared to pure CoOOH nanosheets. The onset potential for the OER was 195 mV for the Mndoped CoOOH nanosheets, while it was around 300 mV for pure CoOOH nanosheets. This means that the Mn-doped CoOOH nanosheets required less energy to initiate the OER. The anodic current of the Mn-doped CoOOH nanosheets was also higher than that of pure CoOOH nanosheets at a low overpotential of 255 mV, where it reached 10 mA cm⁻². This is in contrast to pure CoOOH nanosheets, which required a larger overpotential of around 300 mV to reach the same current density. This data suggests that the Mn-doping effectively boosts the catalytic activity of CoOOH nanosheets for the OER. Furthermore, the Tafel slopes of the Mn-doped CoOOH nanosheets were lower than those of pure CoOOH nanosheets and IrO2, which means that the rate of OER is faster for Mn-doped CoOOH nanosheets. The Tafel slope for 2% Mn-doped CoOOH nanosheets is 38 mV dec⁻¹ and 33 mV dec⁻¹ for 5% Mn-doped CoOOH nanosheets, whereas it is 40 mV dec-1 for pure CoOOH nanosheets and 52 mV dec⁻¹ for IrO₂. Electrochemical double-layer capacitance (Cdl) results also showed that the Mn-confined CoOOH nanosheets possess larger $C_{\rm dl}$ than pure CoOOH nanosheets. The largest $C_{\rm dl}$ of 5% Mn-doped CoOOH nanosheets (8.52 mF cm⁻²) is twice that of pure CoOOH nanosheets (3.38 mF cm⁻²) as obtained from the $C_{\rm dl}$ results, confirming more effective surface area and better charge storage capacity for Mn-doped CoOOH nanosheets. Overall, the data suggest that Mn-doping effectively boosts the OER performance of CoOOH nanosheets by lowering the onset potential, increasing the anodic current at low overpotentials, reducing the Tafel slope, and increasing the electrochemical double-layer capacitance. The exact mechanism of how Mn-doping improves OER performance is not clear, but it is proposed that Mn ions may act as active sites for the OER by participating in the adsorption and dissociation of water molecules, or by changing the electronic properties of CoOOH.

Chen et al.166 used operando techniques, specifically in situ SERS to observe the structural changes of the catalysts under anodic polarization during the OER. This allows for the realtime observation of the catalysts' structural changes while they are actively participating in the OER. The study found that the Co-RT and Ni_{4.1}Co-RT catalysts had similar OER activities, with both catalysts having similar electrochemical properties such as the Tafel slope and exchange current density. However, Raman spectroscopy revealed that the structural changes in the catalysts under anodic polarization were different. For the Co-RT sample, the Raman spectra showed a monotonic decrease in the A1g Raman band intensity at potentials greater than that for the resting state. This decrease in intensity was more pronounced at high oxygen evolution rates (around 4 mA cm $^{-2}$). This structural change continues at relatively high oxygen evolution rates, indicating that the structural transformation of

Co₃O₄ to CoO(OH) is closely related to the OER activity. On the other hand, for the Ni_{4.1}Co-RT catalyst, the Raman spectra were similar to those for the Co-RT samples and showed no significant changes under anodic polarization. This suggests that Ni incorporation did not alter the catalyst structure when the Ni_{4.1}Co-RT catalyst was prepared by the long-time annealing procedure. For the Ni_{5.3}Co-100 sample, the Raman spectra showed a significant structural transformation at 0.2 V. At this potential, the A1g band intensity was attenuated by about 45%, and the E g (487 cm⁻¹) and F 2g (531 cm⁻¹) modes completely disappeared. Simultaneously, two broad peaks at 542 and 600 cm⁻¹, as well as a small shoulder at 490 cm⁻¹, appeared. These new peaks suggest a different local structure for this catalyst at the 0.2 V potential, which may be related to improved OER activity. In summary, the study found that the incorporation of Ni in the Co₃O₄ spinel lattice leads to a structural transformation that may improve OER activity in the Ni_{5.3}Co-100 sample, but this is not confirmed by the study. The Ni_{4.1}Co-RT catalyst showed no significant change in the structure under anodic polarization.167

In a study, Mn doping was found to play an important role in improving the performance of OER catalysts.168 The authors of that study synthesized three samples with different ratios of Mn ions in LDH structure and evaluated their OER performance through a three-electrode configuration in 1 M KOH. The results indicate that the sample with the optimal amount of Mn (Co₁Mn₁ LDH/NF) has the lowest overpotential of 285 mV at 10 mA cm⁻², which is lower than those of other samples and some reported excellent Co-based materials. Additionally, this sample has a current density that is nearly 3 times higher than that of the CoCo LDH/NF sample at 1.8 V vs. RHE, which further illustrates the significant synergistic effect of Mn doping on the improvement of OER performance. Additionally, Mn doping leads to faster reaction kinetics as observed through the Tafel slope which is smaller in the sample with optimal Mn doping. It also increases the stability of the catalyst as observed through the chronoamperometry (i-t) test, where the retention rate of current density for the sample with optimal Mn doping is almost 98% even after 20 hours of testing, and the morphology could also be retained well. Furthermore, the faradaic efficiency was also tested to evaluate the oxygen conversion rates, and the sample with optimal Mn doping is found to have 94% faradaic efficiency, thus exhibiting more stability and efficient energyconversion rate in the material. In summary, Mn doping plays a crucial role in the OER by significantly improving the performance of the catalyst by lowering the overpotential, increasing the current density, and leading to faster reaction kinetics, greater stability, and higher efficiency.

Karmakar and Srivastava studied the electrocatalytic activity of $Co_{1.9}Ni_{0.1}(CO_3)(OH)_2/GP$ and $Co_{0.95}Mn_{0.05}CO_3/GP$ measured for the OER at a scan rate of 5 mV s⁻¹ with *iR*-compensation and the corresponding polarization curves.169 It was found that NiCCH exhibited the lowest overpotential (η) values at all current densities and even outperformed the commercial stateof-the-art RuO2 electrocatalyst. The improvement in OER performance was attributed to the presence of multiple valence oxidation states and increased active metal centers due to the substitution of Co with heterometal atoms (Ni or Mn). Additionally, the larger specific surface area of NiCCH and MnCCH compared to CCH for their superior electrochemical activity was also considered. The 2D rectangular nanoplates of NiCCH and 3D octahedral MnCCH particles could also facilitate electrolyte penetration for better contact with available active sites for the OER. Furthermore, the findings showed that the lowest Tafel slope for NiCCH indicated faster OER kinetics in comparison to RuO2, CCH, and MnCCH. The calculated Rct values suggested a faster electron transfer rate in NiCCH and MnCCH compared to CCH. Overall, the study suggests that transition-metal substitution effectively modified the electronic structure of NiCCH and MnCCH, enhancing their electrical conductivity and OER performance.

The study performed by Zhang et al.93 has shown that the CoIr-0.2 catalyst, which is composed of atomic iridium and Co(OH)2, exhibited excellent OER performance in a neutral electrolyte (1.0 M PBS, pH = 7) and alkaline solution (1.0 M KOH, pH = 14). This is evidenced by its low overpotential (η) required to achieve a current density of 10 mA cm⁻², which is lower than that of commercial IrO2 in both solutions. The Tafel slope for CoIr_{-0.2} is also much smaller than those of other CoIr_{-x} samples and IrO2, indicating that it requires less energy for electrochemical processes. Additionally, the stability of CoIr_{-0.2} in both neutral and alkaline solutions is excellent. X-ray absorption spectra were used to confirm the presence of atomic iridium in the CoIr-0.2 catalyst and to understand the effect of iridium incorporation on the electrocatalytic performance. Overall, the study suggests that the CoIr-0.2 catalyst, composed of atomic iridium and cobalt hydroxide, has great potential for practical applications in the OER.

The study reported by Li et al. 170 demonstrated that the Co(OH)₂/BP nanosheets with a mass ratio of 5:1 showed the best OER activity. The overpotential (η) of Co(OH)₂/BP 5:1 nanosheets at a current density of 10 mA cm⁻² was 276 mV,

which is significantly lower than that of pure Co(OH)₂ (370 mV). Furthermore, the current density (at 320 mV) of Co(OH)₂/BP 5: 1 was 39.6 mA cm $^{-2}$, which is about 20.8 times that of Co(OH)₂. The study also demonstrated that the Tafel slope of Co(OH)2/BP 5:1 is reduced to 57 mV dec⁻¹, compared to 88 mV dec⁻¹ for Co(OH)₂ nanosheets, indicating a sharp increase of current density with increasing the overpotential and facile charge transfer in the nanosheets. Additionally, the study also reported that the Co(OH)₂/BP 5:1 nanosheets possess a much larger ECSA than Co(OH)2, indicating a significantly improved active surface. The study suggests that the addition of black phosphorus (BP) to Co(OH)2 greatly improves the OER activity and stability of the catalyst. It is not specified in the study how exactly BP boosts the OER in this specific case; however, in general, it is believed that BP has high conductivity, good thermal stability, and high surface area which makes it a good candidate for electrocatalysis. BP has a layered structure which makes it a good support for catalytic active materials; it also has a high surface area which allows for more active sites for catalytic reactions. Also, it has high electrical conductivity which can facilitate charge transfer. In addition, its unique properties such as high stability, high surface area, and high conductivity may enhance the electronic interaction between the Co(OH)2 and BP, which could lead to a significant enhancement in the OER activity. It's worth noting that this is only a speculation, and the actual mechanism of how BP boosts the OER of $Co(OH)_2$ in the specific case is not reported in the study.

Dong et al. 171 found that the catalytic effect of Ag-decorated cobalt hydroxide nanosheets is much better than that of pure Co(OH)₂ nanosheets. The amount of Ag in the Co(OH)₂ nanosheets also had a significant impact on the performance, where a certain range of Ag can decrease the overpotential. However, excessive Ag can reduce the catalytic performance of the material by occupying the active sites of Co(OH)2. The optimum catalytic performance was found to be at 10% Ag-

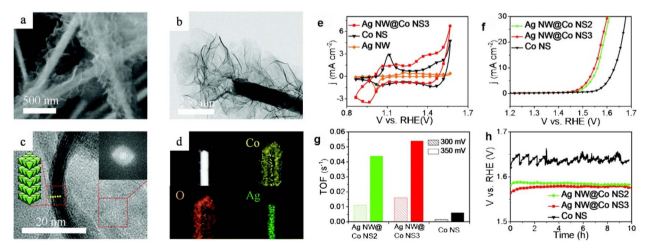


Fig. 11 (a) SEM, (b) TEM and (c) HRTEM focused on the Co NS part and (d) EDX mapping images of the Ag NW@Co NS₃. (e) CVs of the Ag NW@Co NS_3 , pure Co NS and Ag~NW catalysts in an N_2 -saturated 1 M KOH electrolyte at a scan rate of 50 mV s⁻¹. (f) LSVs in an N_2 -saturated 1 M KOH electrolyte at a rotation rate of 1600 rpm and a scan rate of 10 mV s⁻¹, (g) TOFs at overpotentials of 300 and 350 mV and (h) CP tests of the Ag NW@Co NS (2 and 3) and pure Co NS at a current density of 10 mA cm⁻² for 10 h. This figure has been reproduced from ref. 176 with permission from Royal Society of Chemistry, copyright 2016.

decorated Co(OH)2 nanosheets, which had the highest current density of 37.6 mA cm⁻² and the lowest overpotential of 270 mV. It also had a lower Tafel slope of 67 mV dec⁻¹ compared to other Ag-decorated Co(OH)2 nanosheets, indicating a remarkable OER kinetic increment. Overall, the 10% Ag-decorated Co(OH)2 nanosheets showed better catalytic performance than other materials in the study and previous research results.172-175 Kim et al.176 discussed the use of Ag NW@Co NS2 and Ag later@Co NS3 catalysts in the conversion of solar energy to fuels with 10% efficiency. Fig. 11a-d depict the SEM, TEM, HRTEM, and EDX mapping analyses of the sample. These analyses confirm the sheet-like morphology of the samples enveloped by clearly defined Ag nanowires. The EDX mapping demonstrates substantial concentrations of Co, Ag, and O within the sample, validating their presence in significant quantities. The AgO layer on the Ag NW provides a high concentration of active sites for the OER, which increases the catalytic activity of the catalyst. Ag NW also has a well-assembled core-shell structure, which provides more stability to the catalyst layer, and allows for the facile release of O2 bubbles without damage to the catalyst. The study quantifies and compares the OER performance of different catalysts. To explore the surface redox characteristics of the pure Co NS, Ag NW, and Ag NW@Co NS3 catalysts, CV was conducted in a nitrogen-saturated 1 M KOH solution shown in Fig. 11e. The efficiency of the OER was assessed through LSVs depicted in Fig. 11f. Notably, the Ag NWs displayed minimal reactivity within the tested potential range, aside from an anodic peak at approximately 1.6 V (vs. RHE), which was attributed to the oxidation of Ag₂O. TOFs at overpotentials of 300 (1.53 V vs. RHE) and 350 mV (1.58 V vs. RHE), respectively, are illustrated in Fig. 11g. The Ag NW@Co NS2 and NS3 catalysts exhibited TOFs of 0.011 and 0.016 $\rm s^{-1}$, which are 6.9 and 10 times that of the pure Co NS catalyst (0.0016 s⁻¹), respectively, further confirming the higher activity of these catalysts. The stability values evaluated via chronoamperometry of the Ag NW@Co NS₂ (1.56 V) and Ag NW@Co NS₃ (1.55 V) catalysts were found to be lower than that of the pure Co NS (1.63 V) catalyst, indicating their higher efficiency in the OER illustrated in Fig. 11h. The Tafel slope of the Ag NW@Co NS2 and NS3 catalysts (75.8 mV dec⁻¹, 75.4 mV dec⁻¹) was significantly lower than

Liu et al.177 studied the effect of boron (B) and iron (Fe) on the catalysts' OER performance. B is used as a precursor to produce B-α-Co_xFe LDH catalysts. B is found to have an effect on the catalytic activity of B-α-Co_xFe LDH catalysts, with the best activity obtained when using 1 mmol NaTPB. This is likely due to the fact that B can act as a support for the catalytic metal ions and can also affect the electronic and crystal structures of the catalysts, which can improve their electrocatalytic activity. Fe, on the other hand, is one of the metal ions used in the B- α -Co_xFe LDH catalysts. The study found that the B-α-Co_{5.8}Fe LDH had outstanding electrocatalytic activity among the catalysts studied, which suggests that the Fe content plays a crucial role in the catalytic activity. Fe ions may act as active sites or may have a synergistic effect with other metal ions to enhance the catalytic activity. Additionally, Fe can also affect the electronic

that of the physically mixed catalyst (82.5 mV dec⁻¹).

and crystal structures of the catalysts, which can also improve their electrocatalytic activity.

Wang et al.178 synthesized P-doped NiMoO4 parallel arrays anchored on cobalt carbonate hydroxide (NiMoP@CoCH). In this study, the OER activity of electrodes was investigated using corrected LSV curves. The NiMoP@CoCH/CC-2 electrode was found to have a lower overpotential of 267 mV at 40 mA cm $^{-2}$, compared to the other electrodes tested, and also had the highest ECSA. The superior performance of the NiMoP@CoCH/ CC-2 electrode is attributed to its hierarchical heterostructures, which create a "highway" for electrolyte and gas bubble mass transfer, and the incorporation of phosphate dopant and oxygen vacancies, which improves electrical conductivity and increases the number of active defects. Additionally, the electrode's structural and electrochemical stability is enhanced by the growth of NiMoP nanosheets on CoCH nanoarrays. Nevertheless, this electrode has the potential for use in various other fields.179,180

The study performed by He et al. 181 is focused on the electrochemical performance of as-fabricated catalysts towards the OER in 0.1 M KOH electrolyte. The catalysts studied include sol-gel-derived SFM, as-spun SFM nanowires (SFM-NF), bare SFM hollow nanofibers, and Co(OH)2/SFM-NF. The OER activity of these catalysts was initially assessed through LSV polarization curves in a three-electrode system. The sol-gelderived SFM catalyst was used as a benchmark for the study, and its performance was compared to that of the other catalysts. The SFM-NF catalyst was found to have a smaller onset potential of 1.58 V and a larger current density compared to the sol-gel-derived SFM catalyst. This improved OER activity is likely attributed to the larger surface area of the nanofibers. The Co(OH)₂/SFM-NF catalyst was found to have the best OER performance among the catalysts studied, with an onset potential of 1.54 V and the highest current density. This improved performance was attributed to the optimized 10 nm Co(OH)2 decoration on the SFM-NF. The OER current of Co(OH)₂/SFM-NF largely exceeded that of the studied IrO₂ sample beyond 1.60 V, even though the latter had a relatively lower onset potential (1.51 V). The Co(OH)2/SFM-NF sample required a low overpotential of 387 mV to generate 10 mA cm⁻², which is significantly lower than the overpotentials required by IrO2 (447 mV), SFM-NF (468 mV), SFM (548 mV), and Co(OH)₂ (529 mV). The Co(OH)₂/SFM-NF catalyst also exhibited a superior OER activity compared to other reported Fe-based perovskite oxides such as 3D ordered microporous LaFe_{0.8}Co_{0.2}O₃ (460 mV), phosphorus-doped LaFeO_{3-δ} (465 mV), $PrBa_{0.85}Ca_{0.15}Fe_2O_{5-\delta}$ (410 mV), and $SrTi_{0.1}Fe_{0.85}Ni_{0.05}$ -O_{3-δ}/CNT (480 mV). Additionally, Co(OH)₂/SFM-NF had a smaller Tafel slope of 71 mV dec⁻¹ than IrO₂ (88 mV dec⁻¹), SFM-NF (99 mV dec⁻¹), and commercial IrO₂ catalysts (79 mV dec^{-1}). In conclusion, the study found that the $Co(OH)_2/SFM$ -NF catalyst has the best electrochemical performance towards the OER among the catalysts studied, with a smaller onset potential of 1.54 V and the highest current density. The improved performance is attributed to the larger surface area of the nanofibers and the optimized 10 nm Co(OH)₂ decoration on the SFM-NF. The Co(OH)2/SFM-NF catalyst also exhibited

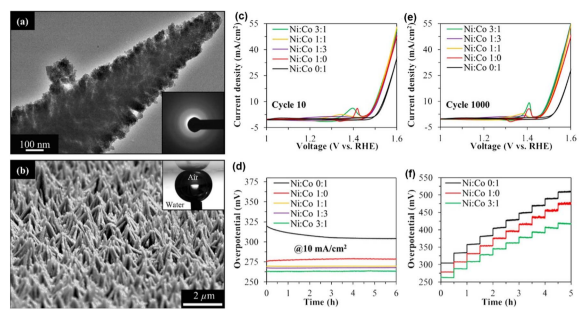


Fig. 12 (a) HRTEM image of a Ni(OH)₂ nanodendrite. Inset shows the selected-area diffraction pattern indicating the amorphous nature of the structure. (b) Tilted cross-sectional view of electrodeposited hierarchical 3-D Ni(OH)2 nanodendrite forests. Inset shows a near spherical air bubble making contact with the deposited surface under water due to ultralow bubble adhesion. (c) Comparison of cyclic voltammograms of nanodendrite (1 min) deposits with varying Ni: Co ratios after 10 stabilizing cycles and (e) after 1000 cycles of accelerated stability testing. All CV curves were recorded at 5 mV s⁻¹. (d) Chronopotentiometry curves for various Ni : Co deposits recorded at j = 10 mA cm⁻² showcase their excellent stability. (f) Chronopotentiometry curves of Ni: Co 3:1 compared to Ni(OH)₂ and Co(OH)₂ deposits recorded at current densities ranging from 10 to 100 mA cm $^{-2}$. Current density values were stepped up by 10 mA cm $^{-2}$ every 0.5 h. This figure has been reproduced from ref. 182 with permission of American Chemical Society, copyright 2017.

a superior OER activity compared to other reported Fe-based perovskite oxides and required a low overpotential of 387 mV to generate 10 mA cm⁻².

4.5 Cobalt based bimetallic hydroxides/layered double hydroxides

4.5.1 Ni-based cobalt hydroxides as OER electrocatalysts. The main focus of the study performed by Balram et al. 182 is on the use of cobalt nickel layered double hydroxide nanodendrites, which were directly grown on a steel substrate, as a catalyst for the OER illustrated in Fig. 12a and b. One of the main advantages of using these nanodendrites is that they are intimately attached to the conductive electrode, which minimizes ohmic losses. Additionally, the lack of binders, conductive carbon additives, and easy electrolyte access, along with their superaerophobic nature, allow for the deposition of a larger amount of material with improved performance. The study found that the Ni : Co 3 : 1 deposit showed the lowest $\eta 10$ mA of ~262 mV, and after 1000 accelerated stability test cycles and 6 hours of chronopotentiometry at j = 10 mA cm⁻², the same samples were immediately subjected to stepped chronopotentiometry from 10 to 100 mA cm⁻² over a 5 h period. Current densities were stepped up by 10 mA cm⁻² increments every 30 minutes. Ni: Co 3:1 outperforms plain Ni(OH)2 and Co(OH)₂ with η10 mA of 262 mV versus 278 and 304 mV for the pure samples respectively. This difference in performance is further amplified at higher current densities with an $\eta 100 \text{ mA}$ value of 417 mV versus 475 and 508 mV respectively (Fig. 12c).

The study also found that despite the prolonged and harsh testing conditions, the deposit showed consistently low overpotentials of \sim 255 mV (Fig. 12d) at j = 10 mA cm⁻² and achieved higher current densities (~15% @ η370 mV) in the OER region compared to the 1 min deposit. However, the performance of the Co(OH)2 sample appears to have deteriorated in the CV curve in Fig. 12b. But the chronopotentiometry carried out at 10 mA cm⁻² after the 1000 cycles of accelerated stability testing shows that the Co(OH)₂ sample returns to the $\eta 10$ mA values exhibited before the harsh cycling process (Fig. 12e). The apparent loss of performance seen in the CV after 1000 cycles may be a function of phase change caused by overcharging during the accelerated stability testing between 1.3 and 1.8 V vs. RHE. Overall, the study provides evidence that the use of nanodendrites as catalysts for the OER can lead to improved performance compared to traditional catalysts, with lower overpotentials and higher current densities. However, further research is needed to fully understand the underlying mechanisms of these nanodendrites and how to optimize their performance.

Prataap et al. 183 designed the porous structure of the NiCo alloy electrode by a single-step electrodeposition method, where hydrogen evolution during the deposition process acted as a dynamic template. This method resulted in the formation of a porous structure with a large surface area, which exposed more electrochemically active sites to the electrolyte. This increased surface area was one of the main reasons for the improved electrocatalytic activity toward oxygen evolution. The

porous NiCo alloy electrode exhibited the highest electrocatalytic activity with the lowest onset potential of 1.47 V and the current density of 10 mA cm⁻² was obtained at 1.53 V. The porous structure of the electrode also enabled more H2O adsorption and accelerated O2 desorption, which contributed to the low OER onset potential and high current density achieved. The porous structure of the electrode also improved the electrode's electrochemical properties. The calculated ECSA value for the porous NiCo alloy electrode was 1.73 cm² and the TOF was 0.0025 s⁻¹, indicating higher catalytic activity. The low solution resistance and charge transfer resistance across the electrode-electrolyte interface, as indicated by low R_s and R_{ct} values, also contributed to the improved electrocatalytic activity of the porous NiCo alloy electrode. In summary, the porous structure of the NiCo alloy electrode played a significant role in improving its electrocatalytic activity towards oxygen evolution in alkaline solutions. The porous structure increased the electrode's surface area, exposed more electrochemically active sites to the electrolyte, enabled more H₂O adsorption, accelerated O₂ desorption, and improved the electrode's stability, electrochemical properties, and catalytic activity. Similarly, Wang et al. 184 synthesized a porous Co_{0.75}Ni_{0.25}(OH)₂ nanosheet catalyst using laser ablation in the liquid method. The catalyst was found to have low overpotentials for the HER and OER and achieved a current density of 10 mA cm⁻² at an external voltage of 1.56 V, which is much lower than those of commercial Ir/C-Pt/C catalysts. The porous structure and the presence of Co³⁺ ions on the pore wall were found to be responsible for the catalyst's high activity. The researchers believe that this laser ablation in the liquid method can be used to create other novel catalysts for energy conversion.185

Wang et al. 186 studied the electrocatalytic OER performance of Co_xNi_{1-x}(OH)(BzO)\$H₂O catalysts with varying Ni/Co ratios. The introduction of a certain amount of Co in Ni(OH)(BzO)\$H₂O greatly enhances the OER activity. The Co_{0.8}Ni_{0.2}(OH)(BzO)\$H₂O catalyst shows the most superior performance among all the samples, with an onset potential of 1.50 V vs. RHE and the smallest overpotential of 319 mV at the current density of 10 mA cm⁻². The introduction of inorganicorganic species and the use of nanobelts have a positive impact on OER activity. It is possible that the inorganic-organic species and nanobelts were used as a way to synthesize the catalysts and their effect on the OER activity was observed as a result. The inorganic-organic species could have played a role in creating catalytically active sites or increasing the active surface area of the catalysts, which would in turn enhance the OER activity. The use of nanobelts may have increased the surface area and improved the electrical conductivity of the catalysts, which would also enhance the OER activity.

CoNi-LDH NSs (nanosheets) were electrodeposited onto a Ni foam by Yoon et al.,187 but the catalytic stability was low (18% degradation) after 10 hours of water electrolysis due to separation from the Ni foam. To improve adhesion, Ni nanocrystals (NCs) were electrodeposited onto the foam to increase the surface area and enhance adhesion. The 0.3/0.2 CoNi-LDH NSs/ Ni NCs/Ni foam exhibited the best catalytic activity and the highest current density (111.5 mA cm⁻² at 1 V). The Ni NCs also

had a positive effect on overpotential and stability, with a current decrease of only 7.4% after 10 hours. Additionally, the Ni NCs did not affect the morphology of the CoNi-LDH NSs. TOF for the CoNi-LDH NSs with the NCs increases from 0.035 to 0.060 s^{-1} in the range from 0.05/0.2 to 0.3/0.2. However, when the Co/Ni ratio is increased further to 0.4/0.2, the TOF decreases to 0.046 s^{-1} .

Keisuke Awaya and Shintaro Ida¹⁸⁸ discussed the electrochemical activity of ultrathin hexagonal nickel-cobalt hydroxide nanosheet films for the OER and ORR. The authors found that a Ni₁Co₄-OH nanosheet film had the lowest overpotential of all the Ni/Co ratios tested and that increasing the layer number of the film also lowered the overpotential. However, the overpotential decreases per layer were larger for 1-4 layers compared to 4-12 layers. The authors also mention that ultrathin films created by stacking nanosheets are ideal for electrochemical reactions and that controlling the layer number of the nanosheets can be useful for investigating the most effective thickness for these reactions.

Wang et al. 189 described the synthesis of a potato chip-like cobalt nickel-layered double hydroxide@polypyrrole-cotton pad (CoNi-LDH@PCPs) by in situ polymerization. This process involves the simultaneous formation of the material and its polymerization. The composite material has a unique potato chip-like structure, which helps to improve the kinetics of the electrode reactions. This material was used as a positive electrode for wearable supercapacitors and it was found to have a high specific capacitance of 387.2 C g⁻¹ at 1 A g⁻¹ and excellent flexibility. As an OER catalyst, the PCP-based electrode had a low overpotential of 350 mV at a current density of 10 mA cm⁻² and a Tafel slope of 58 mV dec⁻¹. The theoretical calculations suggested that the synthesis strategy used in this study can be used to control the formation of hollow CoNi-LDH arrays, which could lead to the development of highperformance electrodes for energy storage and electrochemical catalysis.

Chen et al. 190 presented a new approach for fabricating efficient and low-cost electrocatalysts for hydrogen energy production by fusing NiCo LHS nanowire arrays with ZIF-67 to create three-dimensional flower-like structures on a Ni-Fe foam support. The electrocatalytic performance was improved through simple oxidation, carbonization, sulfurization, and selenization. The optimized S-doped catalyst had the highest electrocatalytic performance, with low overpotentials of 170 and 100 mV for the OER and HER at 10 mA cm⁻² in 1 M KOH, respectively. This was attributed to the strong interfacial coupling between the NiCo-LHSs and ZIF-67 derivatives, as well as the rational electronic structures, dense catalytic active sites, and large specific surface area. The results are promising for the development of next-generation catalytic materials for electrocatalysis and other renewable energy applications.191

Chodankar et al. 192 reported non-noble, precious nanomicro structured Ni₂P₂O₇ microsheets decorated with nickelcobalt hydroxide (Ni-Co-hydroxide) nanoflakes. The designed material exhibited superior catalytic activity towards the OER with a relatively low overpotential of 197 mV at a current density of 10 mA cm⁻² in 1.0 M KOH, coupled with excellent long-term stability (12 h). Outstanding OER activity resulted from the nano-microstructure of the prepared electrode with excellent redox activity in an alkaline electrolyte. The material is stable after 2000 CV cycles at a scan rate of 100 mV s⁻¹ with almost no degradation and shows strong long-term electrochemical stability. It demonstrates high performance with non-noble metal catalysts and a low-cost electrode for oxygen generation. The Ni₂P₂O₇ microsheets in the material serve as the support structure for the Ni-Co-hydroxide nanoflakes. The vertically aligned microsheets provide numerous sites for the electrochemical reactions to take place, as well as a surface to which the Ni-Co-hydroxide nanosheets adhere. The amorphous nature of the Ni-Co-hydroxide with a number of unsaturated metal atoms provides more reactive sites on the catalyst surface, facilitating improved OER. Additionally, the microsheets provide stability to the overall structure, allowing for the longterm stability of the material.

Sun et al. 193 deposited an amorphous boron oxide layer in situ on NiCo-LDH nanosheet arrays through a simple chemical vapor deposition process. This results in a NiCo-LDHs@B₂O₃/ carbon paper (NiCo-LDHs@B2O3/CP) electrode with a low overpotential of 213 mV at a current density of 10 mA cm⁻² in 1 M KOH, which is 123 mV lower than the overpotential of NiCo-LDHs/CP and slightly better than that of the noble metal catalyst RuO₂/CP (230 mV). Additionally, at 50 mA cm⁻² in 1.0 M KOH, the overpotential of NiCo-LDHs@B2O3/CP is still low at 320 mV, which is much better than that of NiCo-LDHs/CP (441 mV) and RuO₂/CP (373 mV). The introduction of amorphous B₂O₃ layers promotes the formation of borate interfaces, which effectively enhances the reactivity of active species and improves the cycling stability of the electrode. This research could inspire the development of low-cost, efficient OER electrocatalysts for energy conversion using NiCo-LDHs@B₂O₃/CP. The B₂O₃ is a known catalyst for the OER. The addition of B₂O₃ to an electrode can increase the rate of the OER and improve the overall efficiency of the process. This is because B2O3 can facilitate the formation of active sites on the electrode surface, where the OER can take place. Additionally, B2O3 can also stabilize the intermediate products of the OER and lower the overpotential required for the reaction to occur, which further enhances the efficiency of the process.

A facile cation-exchange process was developed by Zhou et al.91 to synthesize Fe-doped Ni(OH)2 and Co(OH)2 nanosheets with enriched active sites for enhanced OER. Compared to typical NiFe LDH nanosheets, the Fe-doped Ni(OH)2 nanosheets exhibit higher ECSA and improved surface wettability resulting in significantly enhanced catalytic activity. Specifically, the Fedoped Ni(OH)2 nanosheets require a low overpotential of only 245 mV to reach a current density of 10 mA ${\rm cm}^{-2}$ with a low Tafel slope of 61 mV dec⁻¹, which is greatly decreased as compared with that of NiFe LDH (310 mV and 78 mV dec⁻¹). Additionally, this cation-exchange process can be extended to prepare Fe-doped Co(OH)₂ nanosheets with improved catalytic activity for oxygen evolution. We believe that this novel strategy would shed light on developing high-performance heteroatomdoped catalysts with abundant active sites. Fe is a commonly used dopant in catalysts for the OER because it can significantly

increase the activity of the catalyst. This is due to the fact that Fe can form strong chemical bonds with oxygen, which makes it an effective oxidizing agent. Additionally, Fe has multiple oxidation states, which allows for a greater range of reaction pathways for the OER to occur. Fe dopants are commonly used in a variety of catalysts such as transition metal oxides and hydroxides and have been found to be particularly effective in boosting the OER in non-noble metal catalysts like perovskite oxides and spinel oxides.194

Mixed cobalt and nickel-based LDHs and mixed metal oxides (MMOs) doped with gallium were synthesized by Chakrapani et al. 195 in various compositions by constant pH co-precipitation and calcination. The materials were found to exhibit electrocatalytic activity for the oxygen evolution reaction in an alkaline solution. The highest activity was found for catalysts containing both transition metal cations Co and Ni. However, the effect of calcination was found to depend on the composition of the LDHs. The optimal cation composition of the LDHs was Co_{1.5}-Ni_{0.5}Ga with an overpotential of 382 mV. The highest performance among the MMOs was encountered for the Co_{0.5}Ni_{1.5}Ga composition, reaching a similar overpotential. In particular, gallium has been found to enhance the activity and stability of electrocatalysts used in the OER process. It does this by forming a protective layer on the surface of the catalyst, which helps to prevent the catalyst from degrading or becoming inactive. Additionally, gallium is thought to increase the number of active sites on the catalyst surface, which can further improve the efficiency of the OER process.

4.5.2 Fe-based cobalt hydroxides as OER electrocatalysts. The results of the study by Yang et al. 196 showed that the OER activity of CoAl LDHs decreased with increasing Al content from 15 to 45%, corresponding to an increase of the potential at 10 mA cm $^{-2}$ from 1.63 to 1.68 V vs. RHE, respectively. This decrease in activity was attributed to the dilution of Co sites in CoAl LDHs due to the presence of Al³⁺ ions, which are electrochemically inactive and inhibit the reaction by substituting catalytically active Co sites. However, it is suggested that the improved OER activity of CoFe LDHs compared to CoAl LDHs is due to the presence of Fe3+ ions, which are redox-active and can participate in the redox-hopping-type charge transfer along with Co. The optimal Fe content of 35% was derived for the CoFe LDHs. One possible explanation for the improved OER activity of CoFe LDHs is that the Fe3+ ions act as active sites for the OER, facilitating the transfer of electrons from the Co ions to oxygen molecules. The Fe³⁺ ions may also act as a bridge between Co ions, allowing for faster charge transfer. Additionally, the Fe³⁺ ions may participate in the OER directly by undergoing oxidation to Fe⁴⁺ and donating electrons to oxygen molecules. Another possible explanation for the improved OER activity of CoFe LDHs is that the Fe³⁺ ions enhance the stability of the Co ions in the catalytic active state. It is known that Fe³⁺ ions in the CoFe LDHs can interact with Co²⁺ ions to form Co-Fe pairs, which can stabilize the Co2+ ions against oxidation. This stability may allow more Co²⁺ ions to be available for the OER and improve the overall activity of the catalyst. It is also suggested that the presence of Fe3+ ions in CoFe LDHs may lead to a synergistic interaction between the Fe and Co ions during the

OER. The Fe³⁺ ions may act as a promoter of the OER activity by facilitating the transfer of electrons from the Co ions to oxygen molecules, while the Co ions may act as a stabilizer of the Fe³⁺ ions and prevent them from undergoing unwanted side reactions. It is worth noting that the above explanation is not conclusive, and more research is needed to understand the precise mechanism of how Fe³⁺ ions improve the OER activity of CoFe LDHs. This study is a good starting point for further research, but more in-depth studies, such as Mössbauer spectroscopy, in situ spectroscopy, and density functional theory (DFT) calculations, are needed to fully understand the role of Fe3+ ions in the OER activity of CoFe LDHs.197

The proposed method by Sakita et al. 198 is a one-step, facile, and fast method for obtaining an effective catalyst for the OER. It involves the direct electrodeposition of CoFe alloy thin films covered with LDH in a free-nitrate bath using potentiostatic mode in a continuous or pulsed regime. The catalyst is formed directly on glassy carbon substrates and exhibits a mixed morphology of granular and dendritic CoFe alloy covered with a sponge-like structure of CoFe-LDH containing a chloride (Cl) interlayer. The best-prepared catalyst material showed an overpotential of 286 mV at 10 mA cm⁻², with a Tafel slope of 48 mV dec⁻¹ for the OER, displaying enhanced catalytic activity. The presence of Co and Fe oxidized species at the surface of the composite material and the LDH structure intercalating chloride directly formed on the glassy carbon substrate make it an efficient and cost-effective electrocatalyst for water splitting. The adjustment of the electrosynthesis potential and pulses is crucial to avoid the formation of pure alloy or excessive hydroxides. Additionally, a new approach to preparing a binderfree electrocatalyst has been developed.

A cobalt iron hydroxide (CoFe/NF) electrode was synthesized by Babar et al. 199 through a simple electrodeposition method and showed high catalytic activity and excellent durability in a 1 M KOH electrolyte. The CoFe/NF electrode required overpotentials of 220 mV for the OER and 110 mV for the HER to reach a current density of 10 mA cm⁻². The higher catalytic activity was attributed to synergistic effects between Co and Fe as active sites. The CoFe/NF films were used as bifunctional catalysts for overall water splitting and achieved a current density of 10 mA cm⁻² at 1.64 V and excellent stability for 50 hours. This transition metal hydroxide can be effectively applied for mass production as a highly efficient bifunctional catalyst for overall water splitting. The specific effects may depend on the loading method, loading amount, and other factors. Additionally, other advantages such as effectiveness, high conductivity, and high surface area make Ni foam an attractive substrate for electrocatalysts. 200,201

Guo et al.202 used a simple and efficient electrospinning method to fabricate amorphous cobalt (Co)-iron (Fe) hydroxide on a foamed nickel substrate for the first time. This material showed excellent oxygen-evolution advantages over a typical reference catalyst (RuO2) when tested in an untreated alkaline medium. Additionally, when tested in a three-electrode system in 1 M KOH, the amorphous PVP/CoFe_{1.3} nanofibers had high catalytic activity and stability for the OER, with a low overpotential of 0.267 V at 100 mA cm⁻² and a low Tafel slope of 47.43 mV dec⁻¹. The electrospinning method is considered a feasible and effective synthetic technique for the fabrication of mixed metal hydroxide-based nanofibers with a large surface area, small and uniform grain size, and high morphological uniformity. The study demonstrates the potential of amorphous CoFe hydroxide as a high-efficiency OER catalyst and opens possibilities for large-scale and environmentally friendly production of water splitting. The amorphous nature of the CoFe hydroxide is thought to enhance the OER in several ways. One possible explanation is that the amorphous structure has a higher surface area and a larger number of active sites for the OER to occur, leading to higher catalytic activity. Additionally, the amorphous structure may have a more homogeneous distribution of the Co and Fe atoms, which can promote the synergistic effect between the two elements, further enhancing the OER activity. Furthermore, amorphous materials are often more stable than crystalline materials, which can lead to increased durability of the catalyst over time.203

Defect rich CoFe-based LDHs have been successfully synthesized by Liu et al.204 as a bifunctional, non-noble-metal electrocatalyst for overall water splitting. The delamination and exfoliation process in the DMF-ethanol solvent (CoFe LDH-F) has resulted in an enhancement of both OER and HER activity. The exfoliation process creates more coordinatively unsaturated metals and improves the intrinsic electronic conductivity, which are important factors in water electrolyzer reactions. The CoFe LDH-F catalyst outperforms the commercial IrO2 electrocatalyst in activity and stability for the OER and approaches the performance of Pt for the HER, with a current density of \sim 10 mA cm⁻² at the applied voltage of 1.63 V for the long-term electrolysis test. The performance and costeffectiveness of CoFe LDH-F make it a promising material to replace noble-metal-based catalysts for large-scale water splitting. The defects in the CoFe LDH-F materials play an important role in boosting the OER, which can be further studied to improve the catalytic activity. The defects in the CoFe-LDH structure are believed to enhance its catalytic activity by providing active sites for the OER. In addition, the ultrathin thickness of the CoFe-LDH allows for a high surface area-tovolume ratio, which further increases its catalytic activity.

The investigation found that the Fe sites in the catalyst films were well integrated into the CoOOH structure and that the Fe sites underwent potential-dependent structural changes. Oxidized Fe species play a key role in the OER mechanism on Co(Fe)O_xH_v and the reaction mechanism proceeds through Febased active sites with an intermediate that has Fe in an oxidation state higher than 3+.205

Dai et al.206 described the synthesis of ultrathin carboncoated, nitrogen and iron-modified cobalt carbonate hydroxide needle arrays on carbon cloth (C@NFeCoCH/CC) using a hydrothermal reaction and low-temperature calcination process. The combination of Fe and N modification results in high OER activity of C@NFeCoCH/CC due to the formation of highly active Co/Fe-N-C species on the interface of the carbon layer and needles. The thin carbon layer, N and Fe modification, and continuous conductive carbon cloth allow for fast and stable charge transfer. The optimized C@NFe1Co1CH/CC has

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a low overpotential of 235 mV at 10 mA cm⁻², a mass activity of 681.4 A g⁻¹, and superior stability over 30 hours at 10 mA cm⁻². The introduction of active sites is a promising approach for the application of metal carbonate hydroxides in electrocatalysis. Carbon protection of CoFe LDHs can increase their OER activity by providing a more stable and efficient catalytic surface for the reaction to occur on. Carbon-protected CoFe LDHs have been shown to have higher OER activity and better stability compared to unmodified CoFe LDHs. This is due to the carbon layer protecting the CoFe LDHs from degradation and providing a more efficient catalytic surface for the OER to occur on. Additionally, carbon protection can also increase the electrical conductivity of the CoFe LDHs, which also improves the OER activity.

A stable single atomic ruthenium catalyst was developed by Li et al. 207 through anchoring it on the surface of CoFe LDH. The catalyst was found to have a strong electronic coupling between the Ru and LDH. With a low Ru loading of 0.45 wt%, the catalyst exhibited high activity and stability in the OER, requiring only 198 mV overpotential to drive a current density of 10 mA cm⁻² in an alkaline solution. The activity and stability of the catalyst were attributed to the synergetic electron coupling between the single-atom noble metal and the LDH. These findings have potential implications for the development of cost-effective and high-performance electrocatalysts for energy conversion-related applications.

4.5.3 Other transition metal cobalt-based hydroxides as **OER electrocatalysts.** The most studied cobalt-based hydroxide other than Ni and Fe is the Zn-based Co(OH)2 for the OER. The electrocatalytic OER activities of Co-Zn-LDH were evaluated using a RDE in 0.1 M KOH solution.208 The rotation rate was kept at 2200 rpm to remove the generated oxygen bubbles. The OER activities of bulk ZnCo₂O₄, Zn-CO-LDH nanoparticles, and Co₃O₄ nanopowders were included for comparison. The Zn-CoLDH nanosheet catalyst showed the lowest onset potential and highest current density for the OER among these four catalysts, revealing good OER activity. The comparison of OER activities between Zn-Co-LDH nanosheets and other catalysts was performed from the onset potential, the potential at a current density of 2 mA cm⁻². The Zn-Co-LDH nanosheets presented a lower overpotential (375 mV) at the current density of 2 mA cm⁻² than that of Zn-Co-LDH nanoparticles (460 mV). The powerful effect of the area of Zn-Co-LDH nanosheets on OER activities was found to be consistent with previously reported results. The Tafel plots were also examined. It can be seen that Co₃O₄ had a lower slope of 105 mV per decade, which was lower than that of Zn-Co-LDH nanoparticles (145 mV per dec⁻¹) and ZnCo₂O₄ (157 mV per dec⁻¹). Zn-Co-LDH nanosheets with porous ultra-thin two-dimensional structure exhibited better catalytic performance towards the OER than that of Zn-Co-LDH nanoparticles and Co₃O₄. The use of CoZn-LDH as a catalyst for the OER has several advantages. For example, it is a low-cost and earth-abundant material, making it a more sustainable and environmentally friendly option compared to other catalysts. Additionally, it has been shown to have high activity and stability for the OER, making it a promising material for use in water splitting and other applications.

CoZn-LDH is a type of material that can be used as a catalyst for the OER by using in situ corrosion of zinc foils with cobalt chloride.209 The CoZn-LDH is made up of monocrystalline nanosheets that have a similar lattice orientation and structure coordination which contributes to the thermodynamic stability of the material. The percolated nanosheets are also endowed with hydrophilic functional groups, superaerophobic surfaces, and highly exposed active sites which directly serve as electrodes for electrochemical OER.210 This approach is a simple method for the fabrication of functional nanocomposites for electrical, catalytic, and photochemical applications. A modified hydrothermally synthesized zinc and cobalt-based hydroxyl double salt, Zn_{4-x}Co_xSO₄(OH)₆·0.5H₂O (ZCS), is an efficient electrocatalyst for the OER in alkaline medium.211 The ZCS material was exfoliated for the first time and has unique hexagonal nanoplates. It requires only 370 and 450 mV overpotential to achieve 10 and 100 mA cm⁻² current density, respectively and has a higher mass activity than the commercial RuO2 catalyst at the same overpotential. The ZCS material is cost-effective and environmentally suitable compared to other transition metal-based materials.

The OER activity of CoMn LDH was found to be significantly improved by anodic conditioning (AC).212 AC is a process where the material is exposed to an anodic current density, in this case, J = 10 mA cm⁻², for a certain period. After 3 hours of galvanostatic conditioning at this current density, the overpotential ($@J = 10 \text{ mA cm}^{-2}$) decreased by about 20 mV for CoMn LDH, but only negligibly (3 mV) for Co(OH)₂ + Mn₂O₃. This suggests that the AC process specifically enhances the OER activity of CoMn LDH. Further, it was found that the activity enhancement was durable, with the current density remaining at 16.5 mA cm⁻² at 300 mV for more than 10 hours. The faradaic efficiency, or the efficiency of the catalytic process, was also found to be quantitative during a 5 hour electrolysis experiment. When comparing the activity of CoMn LDH to other OER catalysts, the researchers used TOF which is a measure of the intrinsic activity of a catalyst. The TOF of CoMn LDH at 350 mV (1.05 s⁻¹) was more than 20 times higher than those of Co-Pi, CoOx, MnOx, and their composites (such as mixed Co/Mn spinel), and 9 times higher than that of IrO2. This makes CoMn LDH the most active Co or Mn-based OER catalyst to date. The presence of Mn likely enhances the activity through synergistic effects between the Co and Mn ions, or by providing active sites for the OER process. The researchers mention that the high activity of CoMn LDH is unique and not observed in other Co or Mn-based OER catalysts, which suggests that the combination of Co and Mn ions in this specific CoMn LDH structure plays a crucial role in boosting OER activity.

CoGeO₂(OH)₂ is a cobalt-germanium hydroxide material that has been shown to have potential for use as an electrocatalyst in the OER.213 The results of the LSV measurements showed that the onset of CoGeO₂(OH)₂ occurs before 1.50 V vs. RHE, which means that the electrode begins to produce oxygen at a lower voltage than other cobalt oxides. Additionally, the 10 mA cm⁻² current density occurs at an overpotential of \sim 340 mV, which is lower than the overpotential of 400 mV for most typical bulk cobalt oxides, indicating good OER activity. The mass activation of CoGeO2(OH)2 at 350 mV overpotential is about 49.5 A g⁻¹, which is larger than those of other cobalt compounds, such as Co(OH)₂, Co₃O₄, and CoO. A higher mass activation value indicates that the material is able to produce more current per unit mass and is therefore considered to be more active. This research suggests that increasing the specific surface area of CoGeO₂(OH)₂ may be an efficient route to optimize the OER activity of the material. This is because increasing the surface area would increase the number of active sites available for the OER to occur, leading to a higher overall reaction rate. This has been demonstrated in previous studies on nanostructured Co-based catalysts, where the OER activity was improved by increasing the surface area.

Yang et al.214 reported on the use of freestanding CoVhydroxide nanoneedles as OER catalysts. The best performance was observed with the Co_{0.75}V_{0.25}-HNN catalyst, which had overpotentials of only 266 and 350 mV at current densities of 10 and 100 mA cm⁻², respectively. This is the first report of such low overpotentials for freestanding CoV-hydroxide nanoneedles as OER catalysts and is comparable to most of the reported Co-based and V-based catalysts. The incorporation of V into Co-based hydroxide nanoneedles (NF@Co1-rVr-HNNs) was found to greatly enhance their catalytic activity as OER catalysts. The V incorporation modified the morphology, composition, and electronic structure of the samples, leading to improved OER performance. The V incorporation was found to change the electronic properties of the Co-based hydroxide nanoneedles, resulting in multivalent oxidation reactions, including the V3+ to V⁵⁺ and Co²⁺ to Co⁴⁺ oxidation. This change in electronic properties is thought to be the main reason for the improved OER performance, as it leads to lower overpotentials and higher corresponding currents in the voltage range of 1.25-1.65 V (vs. RHE). The incorporation of V also resulted in an increased active surface area and active catalytic sites, as well as faster reaction kinetics, as indicated by the lower Tafel slopes and higher TOF of the V-incorporating samples. In conclusion, the study shows that the incorporation of V into Co-based hydroxide nanoneedles is an effective strategy to improve their OER performance, by modifying their electronic properties and increasing the active surface area and active catalytic sites.

Ping et al.215 aimed to investigate the use of a novel electrocatalyst, 3DGN/CoAl-NS, for the OER in an aqueous solution of 1.0 M KOH. The 3DGN/CoAl-NS electrocatalyst was prepared by coating CoAl-LDH nanosheets onto 3D graphene. The loading amount of CoAl-NSs on 3DGN was measured to be about 0.05 \pm 0.01 mg cm⁻², using inductively coupled plasma atomic emission spectroscopy. The Tafel slope was found to be 36 mV dec⁻¹ for the 3DGN/CoAl-NS electrocatalyst. This value is lower than the Tafel slope of other reported catalysts, indicating that the 3DGN/CoAl-NS electrocatalyst has a higher reaction rate. The overpotential, which is a measure of the energy required to drive the OER, was found to be 252 mV at a geometrical current density of 10 mA cm⁻². The geometrical current density, which is a measure of the amount of current produced by the catalyst, was found to be 45.37 and 91.74 mA cm⁻² at overpotentials of 300 and 350 mV, respectively. The TOF, which is a measure of the number of OERs that can occur per second, was found to be

0.56 and 1.14 s⁻¹ at overpotentials of 300 and 350 mV, respectively. When compared to state-of-the-art LDH-related OER electrocatalysts, the 3DGN/CoAl-NS electrocatalyst was found to have comparable or even better catalytic activity. Additionally, control experiments were carried out to compare the performance of 3DGN/CoAl-NS with other Al-based LDH nanosheetcoated 3DGN, with 3DGN/CoAl-NS showing the best performance. Overall, the results of this research suggested that the 3DGN/CoAl-NS electrocatalyst has promising potential as a catalyst for the OER in water splitting and other applications. The role of Al in this research was to provide structural support for the electrocatalyst. The Al in the CoAl-LDH nanosheets helped to create a layered structure with a high surface area, which is beneficial for the OER. Additionally, the Al in the CoAl-LDH nanosheets may also contribute to the catalytic activity of the electrocatalyst, by promoting the adsorption of reactant molecules and by facilitating the transfer of electrons during the OER.

Cobalt based ternary hydroxides as OER electrocatalysts

Wang et al.216 evaluated the OER performance of NiCoFe-LDH/ CFC in comparison to bare CFC and NiCo-LDH/CFC. The bare CFC substrate had a very low OER current density and a large onset potential, indicating that the CFC substrate had negligible catalytic activity for the OER process. However, when coated with NiCo-LDH, the material displayed a much-reduced onset potential and an increased current density for the OER. The introduction of Fe to the NiCo-LDH material further reduced the onset potential and enhanced the current density for the OER. The best NiCoFe-LDH/CFC material with 1 hour coprecipitation (CER) time showed a low overpotential of 280 mV, which was 50 mV less than that of the commercial Ir/C based electrode. This demonstrates the superior OER activity of the NiCoFe-LDH/CFC material compared to current commercial OER catalysts. It was found that increasing the CER duration to 3 and 6 hours resulted in a steady or reduced OER activity. This is attributed to the loss of active materials during the CER process, as one mole of Fe(III) replaced 1.5 moles of Co(II) or Ni(II). The overall activity of a material is determined by the intrinsic activity and the amount of active material. In the 3 hour CER sample, the increased intrinsic activity of the electrocatalyst and the loss of active material resulted in a trade-off, while in the 6 hour CER sample, the loss of active material outweighed the benefit of the enhanced intrinsic activity of the material, leading to reduced OER activity. They used Tafel plots to demonstrate the improved kinetics of the Fe-modified material for the OER. The linear range of the Tafel plots of the materials with different Fe content was fitted by the Tafel equation to extract the slope (overpotential). The researchers found that the NiCoFe-LDH/CFC samples with 1 and 3 hours of CER time had a low Tafel slope, indicating improved kinetics for the OER. The lowest Tafel slope was obtained by the 6 hour NiCoFe-LDH/CFC material, indicating an evolution of ratedetermining steps. The Tafel slope of NiCoFe-LDH/CFC was lower than that of NiCo-LDH/CFC, indicating a four-electron transfer determining step, which is a sign of a good

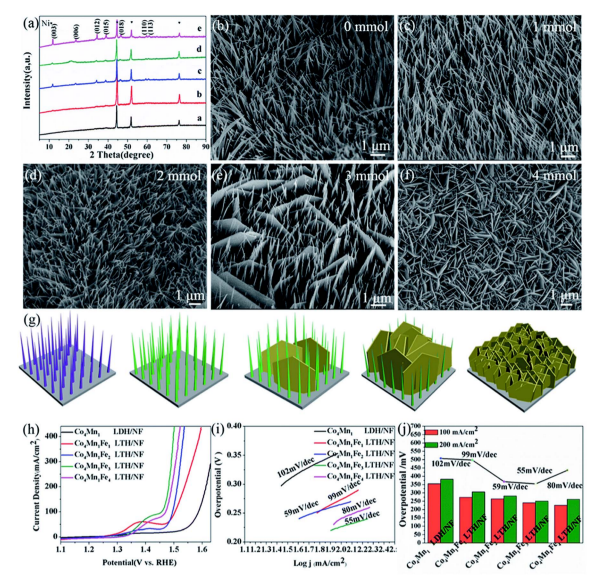


Fig. 13 (a) XRD patterns of a: Co₄Mn₁LDH/NF, b: Co₄Mn₁Fe₁ LTH/NF, c: Co₄Mn₁Fe₂ LTH/NF, d: Co₄Mn₁Fe₃ LTH/NF, and e: Co₄Mn₁Fe₄ LTH/NF samples; SEM images of (b) $Co_4Mn_1 LDH/NF$, (c) $Co_4Mn_1Fe_1 LTH/NF$, (d) $Co_4Mn_1Fe_2 LTH/NF$, (e) $Co_4Mn_1Fe_3 LTH/NF$ and (f) $Co_4Mn_1Fe_4 LTH/NF$ samples; (g) proposed mechanism for the effect of Fe addition on morphology construction; (h) LSV curves, (i) Tafel plots and (j) corresponding contrast of Co₄Mn₁ LDH/NF, Co₄Mn₁Fe₁ LTH/NF, Co₄Mn₁Fe₂ LTH/NF, Co₄Mn₁Fe₃ LTH/NF and Co₄Mn₁Fe₄ LTH/NF. This figure has been reproduced from ref. 229 with permission from Royal Society of Chemistry, copyright 2019.

electrocatalyst. The excellent OER properties of the as-prepared NiCoFe-LDH/CFC are caused by several factors. The in situ CER method preserves the advantages of the NiCoLDH/CFC material, such as an intimate contact between the catalyst and substrate for efficient charge transport, a large surface area, and multiple channels for the diffusion of electrolyte ions. The incorporation of Fe also helps to enhance the material's performance in the OER process by inducing crystal structure disorder and reducing the thickness of the nanosheets, increasing the number of active sites, changing the charge transport determining step, and increasing the conductivity of the electrode.217-228

Guo et al. 229 studied the effect of catalyst morphology on the performance of the OER using a specific material, Co₄Mn₁Fe_y

LTH/NF. XRD patterns of Co₄Mn₁Fe_y LTH/NF products with different Fe levels display similar peaks, matching carbonatecontaining LDH reflections, indicating successful synthesis on the Ni foam substrate illustrated in Fig. 13a. SEM images reveal distinct morphological changes: without Fe, uniform NWs grow (\sim 2–3 µm long, \sim 53 nm in diameter) on the Ni foam (Fig. 13b). With increasing Fe (up to 3 mmol), a transition occurs, leading to a coexistence of nanosheets and NWs forming oriented hybrid arrays, enhancing electron transport (Fig. 13c-e). At 4 mmol Fe, nanowires vanish, leaving only nanosheets, reflecting a significant morphological transformation (Fig. 13f). A schematic diagram illustrating the impact of Fe addition on morphology is provided in Fig. 13g. Various Co₄Mn₁Fe₉ LTH/NF samples featuring different levels of Fe addition were evaluated

for their respective OER performances. Notably, these obtained materials demonstrate suitability for use as self-supporting electrodes without the need for additional coating treatment. Fig. 13h illustrates oxidation peaks in the range of 1.3-1.45 V vs. RHE, typically associated with the transformation of Ni species from Ni foam into Ni(OH)₂ and further into NiOOH via Ni²⁺/Ni³⁺ oxidation. Among the series of electrodes examined, the Co₄-Mn₁Fe₃ LTH/NF stands out with the most exceptional OER performance. It achieves overpotentials of 226 mV and 251 mV at current densities of 100 mA cm⁻² and 200 mA cm⁻², respectively. These overpotentials notably outperform those observed for Co₄Mn₁ LDH/NF (356 mV and 383 mV), Co₄Mn₁Fe₁ LTH/NF (274 mV and 307 mV), Co₄Mn₁Fe₂ LTH/NF (265 mV and 282 mV), and Co₄Mn₁Fe₄ LTH/NF (241 mV and 262 mV) in reaching the corresponding current densities of 100 mA cm⁻² and 200 mA cm⁻² (Fig. 13j). As depicted in Fig. 13j, Co₄Mn₁Fe₃ LTH/NF demonstrates a notably lower Tafel slope of 55 mV dec⁻¹ compared to other electrodes (Fig. 13i). This lower slope signifies highly satisfactory kinetics, wherein a slight increase in overpotential results in a significant rise in current density.²²⁹

Hu et al.230 discussed the use of CoFeV LDH/NF catalysts in 1 M KOH for the OER. The CoFeV LDH/NF catalysts are made by doping cobalt ferrite (CoFe) LDH with V ions. The results show that the CoFeV-0.25/NF catalyst has a lower overpotential and Tafel slope compared to the pristine CoFe LDH/NF, commercial RuO₂/NF, and other CoFeV LDH/NF samples with different amounts of V dopant. The overpotential of the CoFeV-0.25/NF catalyst at a current density of 10 mA cm⁻² is 0.242 V, which is lower than those of CoFe LDH/NF (0.318 V) and commercial RuO₂/NF (0.282 V). This means that the CoFeV_{-0.25}/NF catalyst requires less energy to drive the OER compared to the other catalysts. The Tafel slope of the CoFeV-0.25/NF catalyst is 57 mV dec⁻¹, which is lower than those of CoFeV_{-0.125}/NF (66 mV dec^{-1}), $CoFeV_{-0.5}/NF$ (64 mV dec^{-1}) and CoFe LDH/NF (72 mV dec⁻¹). This confirms the faster OER kinetics and superior catalytic activity of the CoFeV-0.25/NF catalyst. The results showed a slight degradation of the current density over time, indicating the durability of the catalyst. The $C_{\rm dl}$ of the CoFeV 0.25/NF sample is similar to that of the other CoFeV LDH/NF samples, while it is larger than that of the CoFe LDH/NF sample. The intrinsic OER activity of the CoFeV-0.25/NF sample is significantly better than that of the bimetallic CoFe LDH/NF, which demonstrates that the addition of V dopant in CoFe LDH could improve the intrinsic OER activity.

Chen et al.231 presented a comprehensive study on the catalytic performance of CoFeCr (oxy)hydroxides for the OER in alkaline electrolytes. The researchers found that the optimal elemental composition for the Co₅Fe₃Cr₂ (oxy)hydroxides is Co: Fe: Cr = 5:3:2, which exhibited the smallest overpotential $(\eta 10)$ of 232 mV and the highest mass activity of 477.0 A g⁻¹ compared to other CoFe-based catalysts. The amorphous crystal structure of Co₅Fe₃Cr₂ (oxy)hydroxides is also essential for its excellent OER activity as thermally annealing the catalyst at 500 °C resulted in degraded performance. The stability of the catalyst was also evaluated through continuous discharging at 50 mA cm⁻² in 1 M KOH electrolyte and was found to be excellent. Comparison of the catalytic performance of

Co₅Fe₃Cr₂ (oxy)hydroxides to commercial IrO₂ showed that Co₅Fe₃Cr₂ (oxy)hydroxides delivered one of the best catalytic activities among recently reported CoFe-based OER catalysts. The study also compared the catalytic performance normalized by the ECSA of the catalysts, which is important for practical applications. Overall, the research provides a valuable contribution to the field of OER catalysis, and the results suggest that Co₅Fe₃Cr₂ (oxy)hydroxides have the potential to be used as an efficient and stable OER catalyst in alkaline electrolyte.

Wang et al. 232 described a study in which a dendritic CoFeCu ternary alloy was grown on a nickel foam electrode using an electrodeposition method. The dendritic morphology was observed to have formed as a result of the introduction of Cu into the solution, and it was found that the CoFeCu/NF electrode had a large ECSA and fast charge transfer rate. The electrocatalytic activity of the CoFeCu/NF electrode for the OER was found to be excellent, with an overpotential of 202 mV at a current density of 10 mA cm⁻² in 1.0 M KOH. This overpotential is considered low, which is desirable for the OER process. The introduction of Cu significantly improved the OER performance of the CoFeCu alloy compared to the CoFe alloy alone. This improvement was attributed to the enhanced charge transfer and increased exposure of active sites due to the formation of the dendritic morphology. Some transition metals in the CoFeCu alloy had been converted into their hydrogen oxide or oxyhydroxide after electrolysis, which may have contributed to the improved performance. Overall, the results suggested that the dendritic CoFeCu/NF electrode is a promising electrocatalyst for the OER in water splitting, with great potential for practical application. 233,234

Shortcomings and limitations of using cobalt-based layered hydroxides as OER electrocatalysts

Cobalt-layered double hydroxides (Co-LDHs) have been studied widely as a potential catalyst for the oxygen evolution reaction (OER), but there are several shortcomings associated with their

- Stability: Co-LDHs tend to degrade quickly under OER conditions, which limits their long-term effectiveness as a catalyst. Additionally, cobalt ions in the Co-LDH can leach out into the electrolyte, which further reduces the stability of the catalyst.
- · Low activity: finally, Co-LDHs are not as active as other catalysts, such as IrO2 and RuO2, which are widely used in the OER. This can lead to slower reaction kinetics and lower overall performance.
- Limited selectivity: Co-LDHs may not be selective for the OER and may also catalyze the reduction of other species in the electrolyte, such as water to hydrogen.
- Insensitivity to pH: the activity of Co-LDHs for the OER is not strongly affected by pH, which can limit their ability to optimize the reaction conditions for maximum efficiency.
- Low durability: Co-LDHs can be easily damaged by exposure to high temperatures, humidity, and other environmental

factors, which can further limit their stability and longevity as catalysts.

- Cost: Co-LDHs are relatively expensive to produce, which can make them less cost-effective than other catalysts, such as NiO and Ni(OH)₂.
- · Low conductivity: Co-LDHs have low electronic conductivity which reduces their performance as catalysts.

Overall, while cobalt-layered double hydroxides have been studied as a potential catalyst for the oxygen evolution reaction, there are several limitations that need to be addressed before they can be considered as a viable alternative to more established catalysts.

Strategies to improve OER activity on cobalt-based layered hydroxides

Enhancing the oxygen evolution reaction (OER) activity of cobalt-based layered hydroxides encompasses diverse strategic approaches:

- (1) Defect engineering: introducing defects within the crystal structure of cobalt-based layered hydroxides stands as an effective approach. These defects create active sites on the material's surface, enhancing its catalytic activity for the OER.
- (2) Surface modification with synergistic materials: modifying the surface of cobalt-based layered hydroxides using materials like transition metal oxides or conductive polymers generates a synergistic effect. This modification enhances the OER activity of the cobalt-based catalyst.
- (3) Bifunctional catalyst design: employing cobalt-based layered hydroxides in tandem with catalysts such as IrO₂ or RuO₂ produces bifunctional catalysts. This synergistic combination enhances overall OER activity.
- (4) Tailored synthesis methods: utilizing varied synthesis methods to fabricate cobalt-based layered hydroxides with distinct structures (e.g., nanorods, nanosheets) exploits unique properties for improved OER activity.
- (5) Optimization of composition: adjusting the cobalt-totransition metal ion ratio within the layered hydroxide composition optimizes the catalyst's OER activity.
- (6) Enhanced support design: designing suitable supports for cobalt-based hydroxides, like employing carbon-based materials, enhances electronic conductivity and stability, augmenting catalyst performance.
- (7) Stability enhancement in acidic media: addressing the stability of cobalt-based hydroxides in acidic environments involves functionalizing the surface with appropriate groups or applying protective coatings.

Each of these strategies offers promising pathways for bolstering the OER activity of cobalt-based layered hydroxides, providing avenues for further advancement and application in electrocatalysis for various industrial and environmental applications.

The performance of cobalt-based catalysts, in comparison to commonly used OER catalysts like iridium oxide (IrO2) or ruthenium oxide (RuO₂), showcases distinct characteristics. Cobalt-based materials often offer competitive performance

with an OER overpotential in the range of approximately 300-500 mV, while IrO2 and RuO2 tend to exhibit lower overpotentials of around 200-300 mV. However, cobalt-based catalysts present advantages in terms of cost-effectiveness and earth abundance, in contrast to the higher cost and scarcity associated with IrO2 and RuO2. While cobalt-based materials might have slightly higher overpotentials, their comparative affordability and greater availability make them promising alternatives in practical applications despite their slightly inferior performance under specific OER conditions.

Conclusion

In conclusion, the use of cobalt hydroxide and its derivatives as catalysts for the OER in water splitting devices is a promising area of research. Studies have shown that these materials have high OER activity and stability, making them potentially useful in the development of more efficient and cost-effective water splitting systems. Additionally, cobalt is an earth-abundant and relatively inexpensive material, which is another advantage over other catalysts. However, there is still much to be understood about the mechanisms of action of cobalt hydroxide and its derivatives in the OER. Further research is needed to optimize their performance, such as exploring the effect of different synthesis methods, modifying the surface properties of the materials, and studying their stability under different conditions. It is also important to consider the potential environmental and health effects of using cobalt-based materials in large-scale applications. Cobalt is a toxic material and therefore, safety measures should be taken in the handling and disposal of cobalt-containing materials. Cobalt hydroxide and its derivatives have shown promising results as catalysts for the OER, and they have the potential to be a viable alternative to currently used catalysts in water splitting systems. However, more research is needed to fully understand their mechanisms of action and to optimize their performance. Additionally, safety and environmental concerns should be taken into account when considering their use on a large scale.

Conflicts of interest

There are no conflicts to declare.

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