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Superactive NiFe-LDH/graphene nanocomposites as competent catalysts for water splitting reactions

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NiFe-LDH has been recognized as the most efficient and cost-effective material for wider applications in electrocatalytic, photoelectrocatalytic, and photocatalytic water splitting, with supercapacitors and adsorbents, owing to their inimitable physicochemical properties. It is well known that standalone NiFe-LDH executes poor electrical conductivity, sluggish mass transfer, and low activity, which put a question mark on their catalytic efficiency and other applications that require superior electrical conductivity and exciton pair separation efficiency. Most importantly, this constraint creates a hindrance to their superior performance in the area of electrocatalytic and photochemical water splitting. To avoid these shortcomings, the coupled structure of NiFe-LDH/graphene has the potential to reflect properties of both NiFe-LDHs and conductive graphene, which completely overcome the shortcomings of counterparts, ensuring better performance and stability. This review aims to summarize the structural impact of NiFe-LDHs, with the interfacial role of graphene/graphene oxide (GO) by establishing a relationship between their structure and activity. Moreover, the emphasis has been laid on the latest development in NiFe-LDH/GO-based materials, along with attention to synthetic methods targeting the creation of a hierarchal porous nature in the materials with different growth approaches to NiFe-LDH on graphene for applications in electrocatalytic, photoelectrocatalytic, and photocatalytic water splitting activities. The latest research and development in this field using NiFe-LDH/graphene with a sensible intermixing of active sites and conductive framework is explored.

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Introduction

To combat the global climatic scenario, the development of novel technology has proceeded at a sweltering rate that badly affects the zero carbon emission energy demands. As an alternative to fossil fuels, renewable energy technologies have increased enormously in recent decades, offering economically viable strategies in many sectors.¹ However, these technologies have different fundamental aspects in correlation to fossil fuels; consequently, like-for-like substitution is not feasible. Sustainable energy resources, namely wind and solar, are diffuse and intermittent, generating new-fangled challenges for corresponding energy provisions as both time and space demand.^{2,3} Therefore, our generation demands us to develop technologies that could facilitate perpetual energy systems while maintaining stability and reliability in real-time applications. Water splitting is a thermodynamically uphill process and is a green technology that can combat all these energy hurdles with minimal greenhouse gas emissions.4,5 As a product of water splitting, natural H₂ and O₂ gas can be produced from H₂O by the electrocatalytic (EC) method using lowcost electricity, or the photoelectrochemical (PEC) method, or the photocatalytic (PC) method, which might provide the effective use of solar energy.⁶⁻⁸ The EC and PEC water splitting in modern-era electrolyzers are versatile methods for the essential manufacture of green and clean fuels by using sustainable energy resources. In these electrolyzers, the half-cell reaction at the anode is the oxygen evolution reaction (OER, $2H_2O \rightarrow O_2\uparrow + 4H^+ + 4e^-$) and the product O_2 gas, is unconfined.^{9,10} However, extensive overpotential is required in the OER to overcome its sluggish reaction kinetics.¹¹ Similarly, the cathodic half-reaction is the hydrogen evolution reaction (HER, $2H^+ + 2e^- \rightarrow H_2$). In the overall water splitting of H_2O into H_2 and O_2 , gas crossover may be possible in the long run, which raises the issue of safety along with a decrease in energy conversion efficiency due to the back reaction to H₂O in acidic, or OH⁻ ions in alkaline media.¹² A key objective of a good electrocatalyst for electrochemical reactions depends on (i) good electrically active sites for conductivity, (ii) surface area, (iii)

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gas adsorption sites, and (iv) stability.¹³ PEC research strives for the advancement of novel and efficient photoelectrodes. In PEC, solitary photoelectrodes are being utilized for both water electrolysis and solar energy conversion; the photocathode (p-type semiconductor) or photoanode (n-type semiconductor) or both electrodes involve several reaction steps such as (i) the ability to exceptionally harvest light, (ii) easy charge separation and transfer, and (iii) fast OER kinetics.¹⁴ In PEC, band bending occurs at the interface of both electrodes and the electrolyte, which is the dynamic of the force responsible for the separation and transfer of charge carriers. The photoinduced holes transported at the photoanode are responsible for the OER, and electrons at the surface of the photocathode are responsible for the HER. In a heterogeneous photocatalytic water splitting process, photocatalysts are used in powder form and dispersed in polar solvents, and this simple method is more suitable than the PEC method as the economy and reaction conditions using external bias are the main prerequisites in a PEC reaction.^{15,16} The photocatalytic route or lightdriven water splitting to generate molecular H₂ and O₂ using a powdered photocatalyst has tremendous potential for the conversion of solar energy to chemical energy. Under the irradiation of light energy greater than or equal to the energy of the bandgap barrier of a semiconductor, electronic excitation takes place from the valence band (VB) to the conduction band (CB) and leaves behind holes in the VB. The CB electrons can reduce protons to produce H₂, whereas the holes

produced in the VB can oxidize H_2O to generate O_2 gas.¹⁷ An efficient photocatalytic semiconductor mainly depends upon three factors: (i) light absorption capability, (ii) separation and movement of photoinduced charges, (iii) reduction and oxidation reactions occurring on the surface of a catalyst.¹⁸ An inclusive deliberation of the above is the input for addressing the confinement of emerging materials and related energy issues.

Achieving superactive catalysts possessing all these characteristics of high-density active sites with energy generation (H₂ and O_2) by the electrochemical, photochemical or photocatalytic route is a critical research area, and a variety of noblemetal and noble-metal-free catalysts have been explored to fulfill these requirements.9,15 Pt is the best catalyst used for HER but its limitations include stability-related issues and high cost. Similarly, Au and Pd have also been used in electrocatalytic HER, photo/electrocatalytic H₂ evolution applications. Noble metal oxides like RuO2 and IrO2 are established as exceptionally active OER electrocatalysts both in alkaline and acidic media. However, their high cost and limited resources hinder their industrial applications. To overcome these limitations, much attention has been devoted to transition metal (TM) phosphides,¹⁹ nitrides,²⁰ and sulfides²¹ but these oxygenfamily-based TMs possess some drawbacks i.e., corrosion and instability. It is therefore an urgent task to develop catalysts made up of earth-abundant elements of low price, high stability and high efficiency. Recently, layered double hydroxides



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Chemistry. Dr. Parida has made significant contributions toward exploring the chemistry related to structural, morphological, optical, electronic and electrochemical properties of a number of wide bandgap and narrow bandgap semiconductors, doped metal oxides, metal carbides, metal sulphides and nitrides for the effective visible light absorption and charge separation required for efficient photocatalytic water splitting, degradation of pollutants and synthesis of fine chemicals. He has published around 400 papers in high impact factor journals, 7 book chapters, filed 35 patents and supervised 50 Ph.D. students. The important and current relevance of his work is amply reflected in the large number of citations (~18 000) of his work and the H index of 72. (LDHs) containing 1st-row earth-abundant TMs (Ni, Fe, Co, Mn, etc.) have been identified for catalyzing the alkaline OER.²² LDHs involve the isomorphous substitution of a divalent metal cation by a trivalent metal cation, forming brucitelike host layers of $Mg(OH)_2$. The metal cations are octahedrally coordinated by hydroxyl groups forming MO₆ octahedra, where vertexes of the O-H groups are perpendicular to the plane of the host layers that connect to form infinite 2D sheets. LDHs can accommodate charge-compensating organic or inorganic anions and water molecules inside the interlayer in order to compensate for the excess positive charge developed due to the isomorphous substitution.²³ LDHs were first discovered in the form of mineral hydrotalcite as $Mg_6Al_2(CO_3)$ $(OH)_{16}$ ·4 (H_2O) , where the Al³⁺ is substituted for Mg²⁺ in brucite-like layers of Mg(OH)₂, with CO₃²⁻ and water in the interlayer.²⁴⁻²⁶ The chemical formula of LDH is usually represented as $[M(II)_{1-x}M(III)_x(OH)_2]^{x+}[A^{n-}_{x/n}\cdot mH_2O]^{x-}$, where M(II)is divalent and M(III) is a trivalent metal cation. 'A' is the interlayer organic or inorganic anion that compensates for the excessive increase of positive charge in the brucite-like layers. 'n' is the charge on the interlayer anion, m represents the amount of water, 27,28 x is the molar ratio of the trivalent cation, and the density of the surface charge is proportional to the ratio of the trivalent metal cations, x = M(III)/[M(II) + M(III)]. Generally, the compositional range of LDHs resides within 0.2 $\leq x \leq 0.33$. The wide variety of metal cations, metallic ratios M(II)/M(III), and anions associated with LDH are easily exchangeable and create flexibility in the electronic structure and composition.²³ On the atomic scale, diverse metal cations can be used to build stable LDHs; for example: Fe²⁺, Mg²⁺, Ni^{2+} , Co^{2+} , Cu^{2+} , Mn^{2+} , Zn^{2+} , Cd^{2+} , Pd^{2+} and Ca^{2+} are divalent cations and $\mathrm{Co}^{3+},\mathrm{Al}^{3+},\mathrm{Mn}^{3+},\mathrm{Fe}^{3+},\mathrm{Cr}^{3+},\mathrm{Ga}^{3+},\mathrm{V}^{3+}$ and Tb^{3+} are trivalent cations. Tetravalent cations, such as Ti^{4+} (0.61 Å), Zr^{4+} (0.72 Å) and Sn^{4+} (0.69 Å) can be used to build stable LDHs. The most unique property of LDH is that on heat treatment, their hexagonal phase structure is destroyed and they form the mixed metal oxide phase. However, the original LDH structure can be reconstructed by the hydration process of the mixed metal oxide, and these properties are known as the structural memory effect. Additionally, the bulk manufacture of LDHs can be achieved by applying the latest synthetic strategy, followed by their applications in the areas of catalysis, photocatalysis, adsorbents, photochemistry, biomedical science, electrochemistry, magnetisation, and polymerisation.^{27-29,30} The authors, along with other groups, have demonstrated the potential of various pristine or modified binary and ternary LDHs, cation and anion-doped LDH and composites of LDH towards superior energy conversion or energy storage applications. These include using NiFe-LDH,^{27,31-34} MgAl-LDH,²⁸ MgCr-LDH,³⁵ ZnCr-LDH,³⁶⁻³⁸ CoFe-LDH,^{39,40} NiLa-LDH,⁴¹ NiMn-LDH,⁴² CoMn-LDH,⁴³ S-doped NiCoFe-LDH,⁴⁴ Ni/ZnCr-LDH,⁴⁵ MgAlFe-LDH,⁴⁶ etc. Specifically, the authors recently reported various NiFe-LDH-based heterostructured materials for excellent PC water splitting activities.^{27,33,34}

In the early years, Ni and Ni-based oxides were mostly engaged in the OER under basic media.^{47,48} Subsequently,

combinations of Ni with Fe as impurities in Ni(OH)2 were noted to lower the overpotential of OER in Ni-based alkaline batteries.^{49–51} The incorporation of Fe impurities as 0.01% in NiFe has a strong effect in lowering the overpotential of the OER.49-51 Generally, NiFe-based materials such as NiFe alloys,⁵²⁻⁵⁴ oxides,⁵⁴ and (oxy) hydroxides,⁵⁴⁻⁵⁶ are used as earth-abundant catalysts that are effective for the OER under basic media. Active surface areas containing copious amounts of active sites are among the major strategies for fabricating OER electrocatalysts, whereas, in this race, the morphological alteration of NiFe electrocatalysts could fulfill these strategies. Among the TM-based LDHs, NiFe-LDHs is the best-studied material owing to its unique lamellar structures, tunable compositions, easy intercalation, cost-effectiveness, low toxicity, and eco-friendly nature.⁵⁷ A recent study of the OER performances of Mn, Fe, Co and Ni-based (oxy) hydroxides, revealed that the corresponding catalytic performances depend on the OH- $M^{2+\delta}$ bond strength of the order Ni < Co < Fe < Mn.⁵⁸ As a result, extremely strong or weak M-OH bonds can slow down the OER activity. Similarly, a first-principles study revealed that O2 on NiFe-LDHs can be easily generated from OOH* species and the Fe(OH)₃ sites are potentially more active than the Ni (OH)₂ sites. In this aspect, NiFe-LDHs possess inherent catalytic activity towards electrochemical and photoelectrochemical OER in comparison to other TM and noble metal catalysts such as IrO2 and RuO2. 33,34,59-61 From the perspective of a photocatalyst, NiFe-LDH possesses a suitable bandgap (2.2 eV), band edge potentials (CB = -0.01 V, VB = +2.19 V vs. NHE) and intrinsic absorption properties of electronic charge transitions through excited inter-and intra-electronic paths, such as oxo-bridged (Ni²⁺-O-Fe³⁺), d-d transitions in Ni²⁺ cations and ligand to metal charge transfer (LMCT) in $O \rightarrow Ni^{2+}/Fe^{3+}$ which are suitable for PC water splitting activities.^{27,33,34,62} However, the conventional NiFe-LDH with bulky structure suffers from meager photo/electrical conductivity and mass transfer due to deficient active sites and electronic charge transport, and solubility product differences among the metal precursors ((Ni(OH)₂, $K_{\rm sp}$ = 5.5 × 10⁻¹⁶ and Fe(OH)₃, $K_{\rm sp}$ = 2.8 × 10⁻³⁹). Optimal parameters for controlling the structure and composition of LDHs are vital factors for electrochemical and photocatalytic applications.^{33,34,59,60,63,64} Yin et al. reported NiFe-LDH/C that displayed superior OER performances.⁶⁵ These NiFe-LDH/C-coupled systems displayed higher binding energies for Ni 2p and Fe 2p as compared to NiFe-LDH, which was confirmed by the inflections in the local electronic environments of Ni and Fe cations.⁶⁵ The coupling reaction between carbon and NiFe-LDH caused distortions in the electronic environment and formed the Ni/Fe-O-C bond, which resulted in a decrease in the crystal size of NiFe-LDH in the NiFe-LDH/C system.⁶⁶ Hence, conductive substrates or stabilizers like layered carbon materials, such as graphene, could be united to activate the active sites of NiFe-LDH and are together capable of overall water splitting efficiency.67-69

Graphene is packed with carbon monolayers that are arranged atomically in a honeycomb-like structure with superior electron mobility, high conductivity, and an elevated

specific surface area. The conductive carbon lattice of graphene is the anchoring site for nanomaterials to form a variety of functional materials. Importantly, the graphene-based materials, for example, GO, possess unique physicochemical properties and have been widely used in adsorption, electrocatalysis, photoelectrocatalysis and photocatalysis.^{70,71} In addition to the excellent conductivity and large surface area of \sim 2630 m² g⁻¹, the GO matrix contains oxygen functionalities such as hydroxyl, epoxide and carboxylic groups, which can form epoxide-assisted metal oxide networks with NiFe-LDH under ambient conditions. Alternatively, the rich density of the sp² bonded carbon present in rGO is anticipated to maintain the stability of the coupled catalytic system under sluggish OER reactions. The strong metal-rGO bonding interactions, in terms of the overlapping of the dsp and sp² orbitals of rGO with Ni(OH)₂ and the Fe(OH)₃ layer of NiFe-LDH, are responsible for the photocatalytic and electrocatalytic activities.^{72,73} Similarly, NiFe-LDH in combination with heteroatom-doped graphene,⁷⁴ and graphene hydrogel⁷⁵ were reported to exhibit superior water splitting performances. Yet, heteroatom-doped graphene is advantageous because of its chemistry containing defective sites for binding the transition metal cations of LDH.⁷⁴ Hence, NiFe-LDH/graphene could provide equilibration between current collection and the active sites, even altering the electronic environment of NiFe-LDH and, on the whole, increase the kinetics of the charge transfer path.^{65,67}

In general, two-dimensional (2D) nanomaterials give rise to a larger specific surface area and, due to forced dimensional orientation, the electron or charge or ion-pair movement is smoother as compared to bulk materials, and is responsible for the superior electrocatalytic and photocatalytic processes. Motivated along these lines, there are various advanced techniques such as solvothermal/hydrothermal, spin-coating, electrodeposition, ion-exchange, and microwave-assisted synthetic approaches that have been identified for the successful preparation of NiFe-LDH/2D-graphene.13 In most cases, NiFe-LDH/ 2D-graphene can be prepared by a one-step spontaneous coprecipitation method, in which the oxidation of the divalent metal cations to trivalent or higher valence takes place due to the presence of air or O₂ dissolved in solution. A detailed discussion of the synthetic strategy of NiFe-LDH/2D-graphene is provided in the Preparation section below. Moreover, the 3D graphene or GO structure possesses a more axial orientation as compared to the 2D graphene structure, and properties like hierarchical porosity with entangled structure, multidirectional electron mobility, and enhanced conductivity, benefit superior catalytic performances.⁷⁶ The hierarchical porous structure of 3D graphene is a potential substrate that can be jammed with large quantities of catalytically active sites. The combination of NiFe-LDH with 3D graphene (NiFe-LDH/3D graphene) provides a larger surface area, accompanied by electrical conductivity and a hierarchical porous structure for detecting the diffusion of charge carriers in electrocatalytic processes.¹³ In particular, the freeze-drying method has been established as the synthetic process for NiFe-LDH/3D graphene since it generates homogeneous porosity, which causes the

elevated mechanical stability of the interconnected graphene network.⁷⁶ NiFe-LDH/3D graphene has been widely used in various electrocatalytic,^{76–78} photoelectrochemical,^{60,79} and adsorption processes,⁸⁰ but there have been no reports on photocatalytic water splitting activity.

Alternatively, the evolution of the structure and chemical composition can reveal the development of NiFe-LDH/graphene for superior water splitting. In terms of composition, NiFe-LDH/graphene can best be compared with its previous types of derivative materials such as metal oxides,⁸¹ hydroxides,⁸² oxyhydroxides,⁵⁵ and bimetallic alloys, etc.⁵⁵ These LDH-based derivative materials inherit the advantages and inherent properties of a 2D layered structure that exhibits competitive electrocatalytic properties. The structural evolution of NiFe-LDH/graphene can also start from the bulk NiFe-LDH,²⁷ to multifaceted nanostructured morphology including nanosheets (NS),⁸³ hollow microspheres,⁸⁴ and hollow nanoprisms,⁸⁵ etc., with more exposed active sites, which appear to control the agglomeration, and enhance the specific surface area. Alternatively, the hierarchical structure of NiFe-LDH including nanoarrays (NAs),⁸⁶ core-shell arrays,⁸⁷ and pyramidal NAs,⁸⁸ are ideal models with enormous mass and electron transportation properties towards water splitting. Therefore, the structural and compositional transition from the bulk NiFe-LDH to the NiFe-LDH/graphene nanocomposite could demonstrate superior water splitting activities due to the exposure of new active sites and synergistic interactions among components. Fig. 1 provides insight into the latest update in the development of NiFe-LDH/graphene-based materials for the OER and overall water splitting.

Among the vast investigations in these diverse fields of water splitting,^{89–91} NiFe-LDH/2D-graphene and 3D-graphene composites have shown their potential towards activities in the electrocatalytic OER, as an anode material in PEC, and in a few cases as a photocatalyst for H₂ and O₂ evolution.^{22,33,49–56,59} In general, the NiFe-LDH/graphene combination gives deep insight into the structural impact of NiFe-LDHs, with the interfacial function of graphene by establishing a relationship between structure and activity. In recent years, there have been reports work based on either NiFe-LDH or NiFe-LDH/graphene for its oxygen electrochemistry;^{83,92,93} however, much emphasis was placed on the distortion of the lattice structure, synthetic methods, morphology and catalytic studies. Fig. 2 shows the systematic trend in the advancements in NiFe-LDH/graphene-based materials, signifying the most recent synthetic methodologies based on different growth approaches and applications in EC, PEC and PC.

Preparation of NiFe-LDH/graphene nanocatalysts

This section covers the preparation of NiFe-LDH/graphene, based on different growth approaches for superior performance. The growth strategy of NiFe-LDH/graphene is broadly divided into two different approaches: (a) the *in situ* growth of

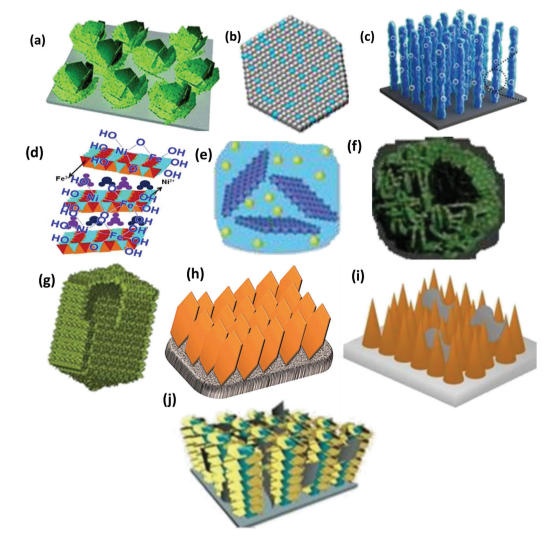


Fig. 1 The structural development of the NiFe-LDH/graphene nanocomposite from the following: (a) metal oxide (e.g., Fe–V@NiO/NF), adapted from ref. 81 with permission from The Royal Society of Chemistry. (b) Hydroxides (e.g. Ni/Ni(OH)₂), adapted with permission from ref. 82, Copyright American Chemical Society. (c) NiFe-oxyhydroxide@NiFe alloy nanowires, adapted from ref. 55 with permission from The Royal Society of Chemistry. (d) Bulk NiFe-LDHs, adapted from ref. 27 with permission from The Royal Society of Chemistry. (e) NiFe-LDH NS, adapted with permission from ref. 83, Copyright American Chemical Society. Nanostructured materials including (f) NiFe-LDH microspheres, adapted with permission from ref. 84, Copyright American Chemical Society. (g) Hollow nanoprisms, reproduced and modified with permission from ref. 85. Copyright 2018, Wiley VCH. (g) NiFe-LDH NS arrays, adapted from ref. 86 with permission from The Royal Society of Chemistry. (h) TiO₂/rGO/NiFe-LDH core-shell NAs, adapted from ref. 87 with permission from The Royal Society of Chemistry. (i) BiVO₄/rGO/NiFe pyramidal NAs, adapted from ref. 88 with permission from The Royal Society of Chemistry.

NiFe LDH on graphene support, and (b) the heterostructure assembly of NiFe LDH/graphene.

(a) The in situ growth of NiFe-LDH on graphene support

The lamellar surface of GO contains functional groups like –COOH and –OH, which offer an appropriate platform for the *in situ* growth of NiFe-LDH.^{94–96} The Ni²⁺ and Fe³⁺ metal cations as the metal precursors of NiFe-LDH possess a strong affinity for, and bond to, the functional groups on the GO surface. Consequently, adsorption and adhesion of metal cations, then nucleation followed by the growth of NiFe-LDH proceeded by the addition of appropriate precipitating agents.

In situ growth can strengthen the linkage between NiFe-LDH and graphene, forming different morphologies.⁹⁷ In the majority of cases, NiFe-LDH/GO originates from a one-step chemical deposition (solvothermal/hydrothermal method).^{33,34,64,98} It is notable that the morphological and structural aspects of NiFe-LDHs fully depend upon the OH⁻ concentration through the nucleation step, and nanometric sizes with tuned morphologies of NiFe-LDH/GO can be obtained by controlling the reaction time and temperature.⁹⁸ Lee *et al.* were the first to report the one-pot growth approach by growing NiFe-LDH on rGO (NiFe/rGO) by a step-wise solvothermal reaction at 120 °C for 18 h, and 160 °C for 2 h

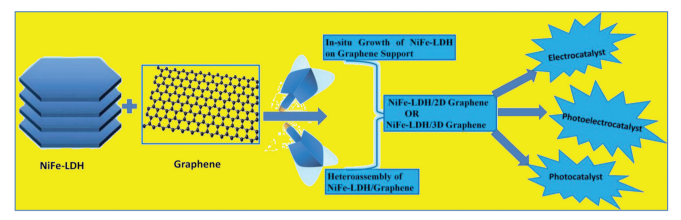


Fig. 2 Scheme of the hierarchical NiFe-LDH/graphene hybrid catalyst with strategies for the synthetic methodology based on two different growth approaches and applications in EC, PEC and PC.

(Fig. 3(a)).⁹⁹ NiFe-LDH was easily crystallized and GO was simultaneously reduced to rGO during the solvothermal reactions. In this approach, NiFe-LDH crystal plates were selectively deposited onto the oxygen functional groups of the rGO layer with higher conductivity and larger surface area. These GO-integrated NiFe-LDH exhibited superior performances in the

electrochemical OER and significantly increased the activity of a Fe₂O₃ photoanode in PEC activity with high stability. Later on, Kang *et al.* explored a related work with the Ni/Fe cation molar ratio of 5:1 and incorporated in GO by using the solvo-thermal technique (Fig. 3(b)).¹⁰⁰ However, Kang's group reported more effective OER activity than that reported by Lee

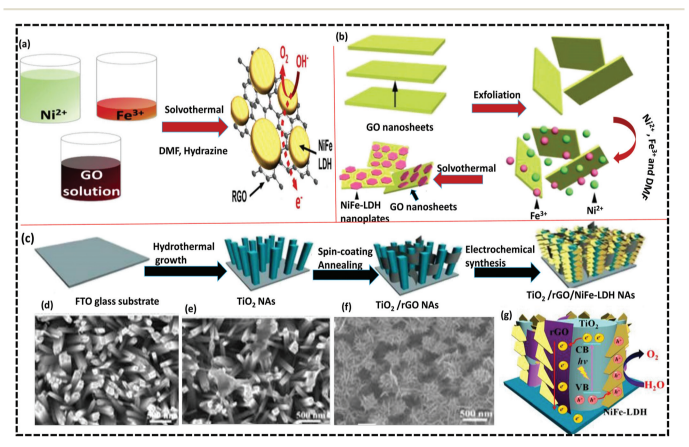


Fig. 3 (a) Illustration of the *in situ* growth of NiFe LDH on the rGO support for the OER. Reproduced with permission.⁹⁹ Copyright 2015, Elsevier. (b) The procedure for the formation of rGO/NiFe-LDH. Reproduced with permission.¹⁰⁰ Copyright 2016, Elsevier. (c) The design process of the TiO₂/ rGO/NiFe-LDH core-shell NAs. (d-f) SEM images of TiO₂, TiO₂/rGO and TiO₂/rGO/NiFe-LDH NAs. (g) The mechanism of PEC water oxidation in TiO₂/rGO/NiFe-LDH core-shell NAs.⁸⁷ Adapted with permission from the Royal Society of Chemistry.

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et al. The electrophoretic method can also be applied to the in situ growth of NiFe-LDH over the GO surface. Generally, a conductive substrate is used for cathodic electrodeposition using Ni²⁺ and Fe³⁺ salts as electrolytes at a constant potential.¹⁰¹ The substrate has a strong impact on the nucleation process. The electrophoretic method is an easy, time-saving, and green method, with a prominent path to building 3D LDH NAs. Interestingly, Shao et al. prepared a NA assembly of a photoanode by spin-coating GO NS over the surface of TiO₂ NAs; the successive electrodeposition of NiFe-LDH nanoplatelets gave rise to the hierarchical core-shell structure of TiO₂/ rGO/NiFe-LDH as shown in Fig. 3(c).⁸⁷ This multistep synthetic approach is reasonably excellent in controlling the composition and morphology by homogeneously anchoring rGO and NiFe-LDH nanoplatelets over the TiO₂ NA surface. The scanning electron microscopy (SEM) image reveals a complete fitting of NiFe-LDH nanoplatelets on the total NAs assembly (Fig. 3(d-f)). These ternary TiO₂/rGO/NiFe-LDH NAs exhibited excellent PEC water splitting activities with a maximum current density of 1.74 mA cm⁻² at 0.6 V and an efficiency of 0.58%, at 0.13 V. The enhanced PEC activities might be attributed to the synergistic result of strong chemical binding interactions among TiO₂, rGO, and NiFe-LDH. Fig. 3(g) displays the mechanism of PEC water oxidation using ternary NAs, which reveals that rGO with a high electron storage tendency, accepts photoinduced electrons of TiO2 and holes are trapped by the NiFe-LDH. Consequently, NiFe-LDH is the active site for water oxidation. Alternatively, the superior electron mobility nature of rGO causes the electrons to rapidly migrate to the PEC current collector. Hence, the role of graphene in this ternary heterostructure is to act like an ideal electron sink, which effectively reduces the bulk recombination and the synergistic effects that enhance the PEC photoconversion efficiency.

The literature study revealed that heteroatom (N, P, S, and B)-doping is an effective approach for increasing the catalytic activities of graphene materials.¹⁰² Generally, melamine and glucose are used as the nitrogen sources for preparing N-doped graphene.¹⁰³ In our reported work, we have also used melamine as a source for doping N in graphene and forming g-C₃N₄ in heterostructured g-C₃N₄/N-rGO/NiFe-LDH.³³ These heteroatoms are called "super-hot spots" (SHS) over the surface of graphene. It was proved that these SHS create topology-induced defects, as well as porosity in graphene. It was also revealed that the SHS contributed to the adsorption of metal cations (Ni^{2+} , Fe^{3+}), followed by the in-plane mesoporous graphene matrix acting as a nanoreactor for the spatially confined growth of NiFe-LDH crystals.^{104,105} Benefitting from the advantages of the porous N-doped graphene framework (NGF), Zhang et al. fabricated NiFe-LDH/NGF in a spatial confinement strategy via a one-step urea-assisted precipitation by using NH₃ as the source of N, and MgO as the template for the deposition of NGF (Fig. 4(a)).¹⁰⁶ The conductive NGF substrate consisting of a mesoporous framework acts as a medium for the in situ growth of ultra nanostructured NiFe-LDH (nNiFe-LDH). It was ascertained that the N-doped graphene induces topological defects that contribute to the adhesion of Ni²⁺ and Fe³⁺ metal cations and promote nucleation and thereby, the mesoporous

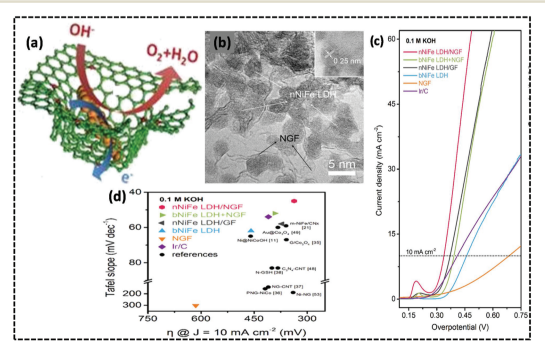


Fig. 4 (a) Diagrammatic representation of the spatially confined hybrid structure of nNiFe-LDH/NGF. (b) HRTEM image of nNiFe LDH/NGF hybrids displaying the homogeneous deposition of nNiFe-LDHs in the NGF matrix. (c) LSV curves of the nNiFe-LDH/NGF series and related materials in 0.1 M KOH electrolyte. (d) The subsequent Tafel plots. Reproduced and modified with permission from ref. 106. Copyright 2015, Wiley VCH.

in-plane graphene acts as a nanoreactor to perform the spatially confined growth and homogeneous dispersion of nNiFe-LDH on the support of the mesoporous graphene framework.¹⁰² The confined growth of nNiFe-LDH within the mesopores of NGF leads to NS of size less than 10 nm (Fig. 4(b)). This process signifies the controlled hybridization of the active phases of nanometric NiFe-LDH into the conductive NGF substrate to form the nNiFe-LDH/NGF hybrid with strong interfacial interactions, which fully exposes the electrochemically active surface area for to facilitate the OER activities (Fig. 4(c)). It is the NGF that acts as a conducting substrate with a high potential for superior charge transport in electrocatalytic reactions. The X-ray photoelectron spectroscopy (XPS) data of nNiFe-LDH/NGF reveals a higher binding energy shift of Ni 2p (ca. 0.7 eV), signifying the linkage of the C-N…Ni-O bond and strong local electronic coupling between NiFe-LDHs and graphene will occur. Consequently, the intrinsic catalytic activities of NiFe-LDHs are tuned with the exposure of active phases like γ -Ni_{1-r}Fe_rOOH or γ -NiOOH, which promotes intermediate adsorption for the OER performance.55,56 Accordingly, the hierarchical structure of the NiFe-LDH/NGF displayed a more kinetically active OER catalyst than Ir/C and other reported materials, with extensive durability in alkaline medium. Important OER parameters include a lower Tafel slope (45 mV dec^{-1} , Fig. 4(d)), small overpotential (337 mV) at boosting the current density of 10 mA cm⁻² in 0.10 M KOH. Overall, the better OER activity of nNiFe-LDH/NGF could be due to the synergistic effect of LDH and NGF with unique structural features due to the involvement of the conductive mesoporous NGF, which facilitates electron and mass transfer through

multiple electronic paths during the OER. Most importantly, the spatially confined growth of nanosized NiFe-LDHs forming a strong hybrid with graphene and generating open coordinate sites and lengthening the edges with exposed surface active sites is responsible for the superior OER performance.

In addition to the above advantages including the high conductivity of the N-doped graphene, another approach is the design of various hierarchical porous structures (such as sphere mesopores, wrinkled mesopores, and strutted macropores) by using a surfactant to improve the potential in the OER.¹⁰⁷ By utilizing the benefit of the porosity of a material for superior catalytic activities, Zhan et al. reported mesoporous NiFe-LDH/NrGO nanospheres by using protonated g-C₃N₄ NS (p-CNNS) as a rich source for N-doping and sodium dodecyl sulfonate (SDS) as a structure-directing agent in a one-pot solvothermal template method (Fig. 5(a)).⁷² The creation of the hierarchical mesoporous/nanospheres could be mainly credited to the reverse micelle effect. The overall catalytic activities of NiFe-LDH/N-rGO in the ORR and OER confirmed its tolerance for long-term durability. The creation of 3D porous nanospheres in NiFe-LDH/N-rGO bestowed it with superior activity due to the increase in the number of active sites and faster mass transfer. Graphene aerogels (GA) also have the advantages of multidimensional electronic channeling and tangled hierarchical porous block structure (mesoporous, and microporous) with high specific surface area, appropriate for the growth of NiFe-LDH.¹⁰⁷ GA also possesses high electrical conductivity and better mechanical properties, which facilitate rapid mass transport of opposite charges, and easy catalyst

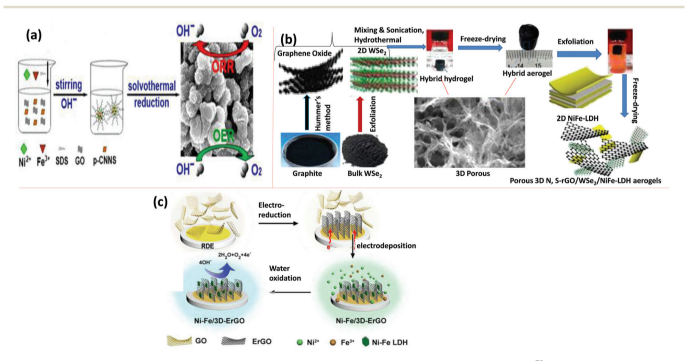


Fig. 5 (a) Schematic diagram for the synthesis of the NiFe-LDH/NrGO catalyst. Reproduced with permission.⁷² Copyright 2017, Elsevier. (b) Synthetic steps of N,S-rGO/WSe₂/NiFe-LDH. Adapted with permission from ref. 79 @ Copyright 2017, American Chemical Society. (c) The design process and application of the NiFe/3D-ErGO electrode.⁷⁸ Adapted with permission from the Royal Society of Chemistry.

recyclability is another merit. Xu et al. prepared porous 3D N, S-rGO/WSe₂/NiFe-LDH aerogels by assembling NiFe-LDH NS with GA via a simple hydrothermal and electrostatic selfassembly technique as shown in Fig. 5(b).⁷⁹ Along these lines, scientists are also capable of designing the one-step synthesis of NiFe-LDH/GA. Despite the exciting experiments in the design of various catalysts for use in electrocatalytic and PEC applications, the higher overpotential, durability and specific current density are the key factors for determining the catalyst potential at specific reactions. The in situ deposition of material over a 3D conductive substrate to design a catalyst or directly depositing the catalyst over a film or a conductive substrate is expected to be more reactive than directly dripping materials on GC electrodes during the activity test. In nanocomposites, the buffering of the electrolyte in a 3D hierarchically macroporous structure increases the ion mobility and utilizes the active species more frequently to facilitate the diffusion of gas bubbles. The most efficient 3D conductive substrates for growing materials are Ni foam,¹⁰⁷ graphene hydrogels,¹⁰⁸ CFP,¹⁰⁹ and the Au-rotating disk electrode (Au-RDE),⁷⁸ or some other metal foil/mesh with carbonaceous substrate were also taken in the precursor metal salt solution, so that the corresponding NiFe-LDH could be developed on it. Shi et al. designed a novel electrode by using NiFe-LDH and 3D electrochemically active rGO (NiFe-LDH/ 3DErGO), supported on Au-RDE by electrochemical reduction of dispersed GO sheets in aqueous electrolyte, and next electro-

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Ultrasonic

GO

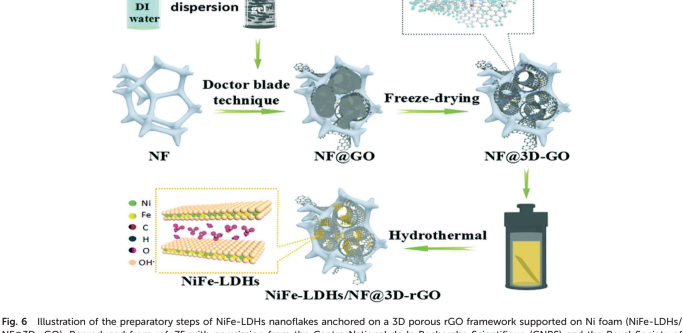
chemical deposition of NiFe-LDH nanoplates on the 3D-ErGO substrate (Fig. 5(c)).⁷⁸

Besides the effects of the heteroatom-doped porous graphene, graphene aerogel and free-standing electrodes, the coupling of NiFe-LDH with GO hydrogel is another novel strategy for boosting the electrocatalytic OER performances. Gu et al. used the GO hydrogel matrix supported over Ni foam by the doctor blade method, which was subsequently freezedried, to produce the 3D-porous GO matrix (Fig. 6).75 NiFe-LDH nanoflakes were attached to the 3D-GO matrix followed by the in situ reduction in a hydrothermal reaction to form 3Dporous rGO/NiFe-LDH supported on Ni foam (NiFe-LDHs/ NF@3D-rGO). The material prepared in this method had a high degree of porosity along with good mechanical strength. The designed 3D-rGO/NiFe-LDH demonstrated excellent OER performance under alkaline medium at the current density of 20 mA cm⁻², with a small Tafel slope of 57 mV dec⁻¹, and overpotential of 170 mV.

(b) Heteroassembly of NiFe-LDH/graphene

3D porous networks structure

The heterostructure is the interface formed due to band alignment among two or more semiconductors with unequal band structure, which possesses intrinsic properties of the countercomponents with considerably enhanced performances. In particular, the heterostructure coupling of NiFe-LDH/graphene synergistically played a vital role in manipulating the band



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NF@3D-rGO). Reproduced from ref. 75 with permission from the Centre National de la Recherche Scientifique (CNRS) and the Royal Society of Chemistry.

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alignment and electronic band structure and facilitated the excessive electron transfer through an interconnected electron highway that promoted water splitting reactions.¹¹⁰ In recent vears, exfoliation has been recognized to be a simple method for generating thin layer NS with a theoretically calculated higher specific surface area of 1000 m² g⁻¹, which is associated with huge quantities of exposed active sites for enhanced catalytic efficiency. Solid NiFe-LDH, possesses strong interplane chemical bonds but weaker out-of-plane van der Waals forces.¹¹¹ NiFe-LDH can be exfoliated and self-assembled with other layered materials like graphene, such as g-C₃N₄ and MoS₂, to form heterostructured materials. We have also reported different heterostructured materials of NiFe-LDH by the exfoliation strategy, composited with other layered materials such as in g-C₃N₄/NiFe-LDH,²⁷ Ag@Ag₃PO₄/g-C₃N₄/ NiFe-LDH,⁶² g-C₃N₄/N-rGO/NiFe-LDH,³³ and MoS₂/NiFe-LDH.³⁴ The simplistic exfoliation strategy of NiFe-LDHs can alter the electronic band structure and generate more active sites with higher conductivity, thus radically increasing the potential catalytic activity and stability. In the past years, various methods for exfoliating a material have been cited in the literature, such as liquid-phase, plasma-induced, and solid-phase exfoliation, etc.7 Our recent work also demonstrated the caliber of the exfoliation technique in the design of the ternary NiFe-LDH/N-rGO/g-C₃N₄ heterostructure. This ternary heterostructure was synthesized by multistep calcination-electrostatic self-assembly and hydrothermal routes.³³ The HRTEM image manifests the intercalation of N-rGO at the interface of the NiFe-LDH/g-C₃N₄ nanohybrid (Fig. 7).³³ The extraordinary photocatalytic activity of this ternary heterostructure towards PC H₂ and O₂ evolution along with effectiveness towards the degradation of RhB dye and phenol might be due to the synergistic electrostatic coupling interactions of the constituent semiconductors and the unique role of N-rGO as a mediator or electron transport bridge in the effective charge separation through the *Z*-scheme mechanistic pathway.

As stated above, the electrostatic self-assembly method is a viable technique generally used for the construction of the NiFe-LDH/graphene heterostructure. It utilizes the opposite surface charge of two positively and negatively charged materials bonded through electrostatic forces of interaction. The aqueous suspension of graphene is negatively charged, while the positively charged host layer of NiFe-LDHs is built up with an orderly arrangement of metal cations and these are the fundamental properties for designing the heteroassembly of NiFe-LDH/graphene via electrostatic self-assembly.33 In 2014, Yang and Long et al. reported the strongly coupled NiFe-LDH/GO heterostructure by the anion (CO₃²⁻ or Cl⁻) intercalation method to enlarge the interlayer space of NiFe-LDH, followed by exfoliation and self-assembly with GO sheets.¹¹² The opposite stacking of NiFe-LDH and GO NS resulted in enormous exposure of the large active surface with higher conductivity (Fig. 8(a)). Further progress was acquired by the reduction of the heteroassembled GO to rGO (named FeNi-GO/ rGO LDH) for the high electrocatalytic OER performance. The SEM morphologies of FeNi-Cl LDH and FeNi-GO LDHs revealed rough surfaces and smaller platelets as compared to FeNi-CO₃ LDH (Fig. 8(b-d)). The reduction of GO in FeNi-rGO LDH was further confirmed by the XPS spectra and XRD patterns. It is worth mentioning that the as-synthesized FeNi-GO

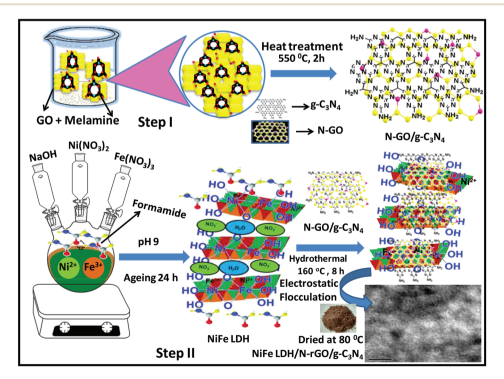


Fig. 7 Synthetic methods of the NiFe-LDH/N-rGO/g-C₃N₄ nanocomposite. Adapted with permission from ref. 33, Copyright 2019, Nature.

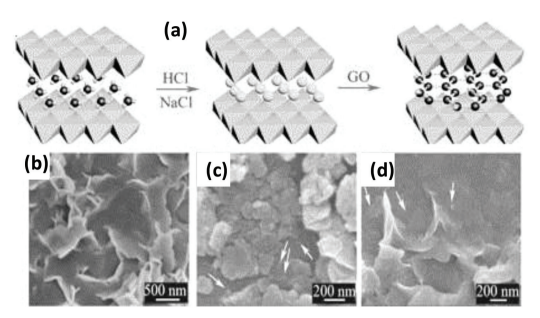


Fig. 8 (a) Fabrication process of the NiFe-LDH/GO hybrids. (b-d) SEM images of NiFe-CO₃ LDHs, NiFe-Cl LDHs, and NiFe-LDH/GO hybrids. Reproduced and modified with permission from ref. 112. Copyright 2014, Wiley VCH.

LDH showed an incremental basal spacing of 1.1 nm, more than twice that of FeNi-CO₃ LDH (0.75 nm), which shows the intercalation of GO into the FeNi-LDH. The extended basal spacings of FeNi-LDH permit the diffusion and association of the electrolyte and reactants. The synergistic strong interaction between the stacked rGO layers hybridized with the interlayer of FeNi-LDH by the layer-by-layer assembly facilitated charge transport through the exposed active sites of FeNi-LDH on the conductive rGO layers and thereby led to the favorable OER performance.

Hybrid structures consisting of catalytically active species and conductive supports are the best strategy for improving the electrocatalytic performance. By this approach, constructing a 2D superlattice structure for achieving targeted activity could be far better than preparing simple composites in which lower intimate contact is possible between the active edges and the conductive surfaces, which lead to less migration of electrons or less ion transfer. Sasaki et al. fabricated the NiFe-LDH/GO superlattice by a facile delamination strategy (Fig. 9(a)).⁸³ In this approach, positively charged NiFe-LDH NS were produced by exfoliating the bulk counterpart in formamide. In the meantime, negatively charged GO NS were synthesized by exfoliating bulk graphite via a modified Hummers' method.¹¹³ Afterwards, the superlattice NiFe-LDH/rGO nanostructure was realized by the electrostatic flocculation between the electrocatalytic active sites of NiFe-LDH and the conductive GO/rGO surface through electrostatic interaction forces. In this process, a synergistic effect from the layer-by-layer assembly between NiFe-LDH and GO with interfacial hybridization is responsible for the formation of the superlattice structure. Accordingly, the NiFe-LDH/rGO superlattice has the advantages of the redox potential of NiFe-LDH and the conductive nature of graphene, which facilitate the achievement of a small Tafel

slope of 40 mV dec⁻¹ at the overpotential of 0.217 V, with a current density of 10 mA cm⁻², and allow this material to function as a NiFe-based electrocatalyst. Similarly, the CoAl-LDH/GO superlattice was prepared by the layer-by-layer assembly and was used as a potential electrode material for supercapacitor devices.¹¹⁴ In contrast to other superlattice structures, $MoS_2/graphene$ was prepared by utilizing the advantages of both the metallic 1 T phase MoS_2 NS and graphene NS, which showed excellent properties in the HER and sodium storage.¹¹⁵

In addition to the aforementioned superlattice heterostructure, the 3D array-like heterostructure is considered to be an emerging material that fully exposes all active sites, which is most effective for energy conversion. However, the stability of the unilamellar NiFe-LDH NS in the 3D array-like heterostructure is still challenging. The stability and directional growth evolution of monolayer LDH NS on GO (horizontal spreading or vertical alignment) can be well controlled by the solid-state exfoliation strategy. By the use of a solid-phase exfoliating material (SPEM) such as a polyhydroxy material, Shen et al. exfoliated NiFe-LDH and GO by using the solid-phase exfoliation strategy to form the 3D array-like NiFe-LDH/GO heterostructure (Fig. 9(b)).⁷³ This process revealed the dynamic growth evolution of NiFe-LDH NS on GO NS (horizontal spreading or vertical alignment or both) and provided information about the fully exposed active sites for interfacial interactions. The NiFe-LDH/RGO heterostructure with the vertical growth of NiFe-LDH NS displayed superior electrocatalytic OER activity with a small Tafel slope of 49 mV dec⁻¹ and overpotential of 273 mV at exchange current density of 30 mA cm⁻² under basic medium. The superior electrocatalytic OER activity of NiFe-LDH/RGO could be accredited to the strong interfacial coupling interactions between NiFe-LDH and RGO NS, as well as the synergistic effect. All these strategies mentioned in this

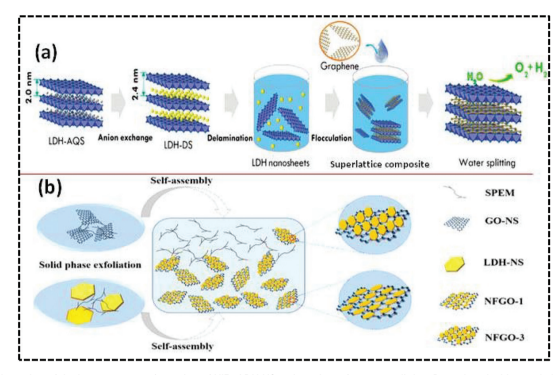


Fig. 9 (a) Illustration of the heterostructure formation of NiFe-LDH NS and graphene for water splitting. Reproduced with permission from ref. 83, Copyright 2015, American Chemical Society. (b) Synthetic steps of NiFe-LDH/GO nanohybrids by using the solid-phase exfoliation–liquid-phase assembly strategy. Reproduced with permission from ref. 73, Copyright 2019, American Chemical Society.

section, including the growth of highly intrinsic LDHs on conductive graphene or the heteroassembly, provide effective methods with morphological tuning, directional growth, contact sites, reaction conditions, and reaction parameters, which are important parameters that determine the formation of NiFe-LDH/graphene hybrid for diverse performances.

Applications of NiFe-LDH/graphene

The ordered structure and tunable compositions of NiFe-LDHs furnished innovative ideas for the design of extremely competent catalysts for applications in EC, PEC, and PC.^{27,87,116,117} GO is a carbonaceous material responsible for enhancing electron conductivity, as well as providing an enormous surface area for the homogeneous growth of NiFe-LDHs. In this section, we mainly emphasize the applications of NiFe-LDHs/graphene in the electrocatalytic OER, and the photoelectrochemical and photocatalytic H₂ and O₂ evolution reactions.

OER catalysis

Water electrolysis is the most exciting scientific achievement due to the simplistic path for electrochemically breaking of H_2O into clean H_2 and O_2 . The only starting material is H_2O and low-cost electricity from solar, wind and hydropower or photovoltaics.^{4–11} Generally, water electrolysis is carried out in a quartz cell consisting of a cathode and anode in which the HER and OER take place.¹¹⁸ The HER is a simple electrochemical reaction that requires a lower overpotential to drive the kinetics of the reduction reactions at the surfaces of many metals. The OER is the most important half-cell reaction for water electrolysis. It involves four proton-coupled electron transfer pathways (4e⁻ transfer and 4H⁺ removal from H₂O), with the sluggish kinetics and high overpotential that decrease the efficiency of the overall water splitting. It is preferred that molecular O₂ be generated from the metal oxide surface rather than neat metal, and therefore its mechanism is guite elusive and confined to specific systems. The majority of metals are transformed into hydroxides or oxides throughout an OER process; however, stability in a harsh acidic medium remains challenging, excluding IrO2 and RuO2. LDHs systems are very sensitive to acidic media and hence, the OER of LDHs is normally conducted in alkaline media. Usually, the protons couple in four steps and the electron transfer routes in heterogeneous electrocatalysis in alkaline media occur as follows:

$$[M] + OH^- \leftrightarrow [M - OH] + e^-$$
 (1.1)

$$[M - OH]_{ads} + OH^- \leftrightarrow [M - O] + H_2O + e^-$$
 (1.2)

$$[M - O]_{ads} + OH^- \leftrightarrow [M - OOH] + e^-$$
 (1.3)

$$[M - OOH]_{ads} + OH^- \leftrightarrow [M - OO] + H_2O$$
 (1.4)

$$[M - OO]_{ads} + OH^- \leftrightarrow M + O_2 + e^-$$
 (1.5)

Unlike the HER, a low overpotential of 10 mA cm^{-2} , a small Tafel slope, higher reaction dynamics to facilitate the multi-

step electron transfer, and easily available metal-containing active sites are the main criteria that determine the potential of an electrocatalyst to satisfy the OER.¹¹⁹ The Tafel slope is very much dependent on the applied potentials. At a higher positive potential and higher Tafel slope, the electrode becomes positive due to the availability of OH⁻. Consequently, the rate-determining steps of the OER involve the formation of M-O bonds or the adsorption of OH⁻. Similarly, at a smaller Tafel slope, the rate-determining step involves the formation of M-O and M-O-O bonds, respectively. However, the OER suffers from intrinsically sluggish kinetics and larger overpotentials. To date, noble metal catalysts such as RuO₂ and IrO₂ have been used in the OER, but they are costly and suffer from instability issues. Current research has explored the potential of earth-abundant transition metals and their derived compounds to replace these noble metals for the OER.¹²⁰ Since Ni²⁺ and Fe³⁺ are the most suitable transition metal cations for the OER, NiFe alloys have been reported to exhibit the electrocatalytic OER.⁵²⁻⁵⁴ Alternatively, the NiFe-LDHs catalysts are the most appropriate candidates for the electrocatalytic OER due to their tunable structure, composition and stability in the electrochemical OER. Candelaria et al. explored the impact of Fe in controlling the electrocatalytic activities of NiFe nanoparticles (NP).¹²¹ Bimetallic FeNi NP evolved with a core-shelllike structure, consisting of $Fe(OH)_3$ or α -Fe₂O₃ as the core and $Ni(OH)_2$ integrated with metallic Fe as the shell. Monometallic Ni NP contains both β -Ni(OH)₂ and α -Ni(OH)₂ with a consider-

able quantity of metallic Ni NP. Interestingly, bimetallic FeNi NP was found to have a significantly higher OER activity than the monometallic Ni and Fe due to the Fe doping in $Ni(OH)_2/$ NiOOH. Zhang et al. subsequently reported a highly competent NiFe film that exhibited excellent OER activities;¹²² specifically, Fe³⁺ was the main modulator for controlling the valence states of Ni in the electrocatalytic processes. The surface defects created by the doping of Fe³⁺ helped in the charge transfer and migration to coordinate the catalyst. Further, Swierk et al. systematically increased the presence of active sites in $Fe_xNi_{1-x}OOH$, by incorporating Fe^{3+} into NiOOH.¹²³ They found that the OER activity of Fe_rNi_{1-r}OOH was faster as compared to FeOOH or NiOOH by using the impedance and activation energy measurement techniques. Also, the faradaic resistance of the monometallic catalyst was found to be 2-fold higher as compared to $Fe_xNi_{1-x}OOH$, whereas the activation energy was 3-fold higher. Besides, NiFe-LDH was found to contain Ni²⁺ and Fe³⁺ cations at low potential.¹²⁴ At an elevated potential, Ni²⁺ is oxidized to Ni³⁺ with the formation of the γ phase of Ni_{1-x}Fe_xOOH, and Fe³⁺ remains unaltered (Fig. 10(a)).¹²⁴ DFT+U studies also verified the phase transformation of NiFe-LDH to Fe-doped γ -NiOOH during the OER, which caused the shifting of the adsorption energies of the oxygen intermediates. Mostly, Fe³⁺ content increases the OER activity of NiFe-LDH by lowering the overpotential of γ-Ni_{1-x}Fe_xOOH as compared to Ni³⁺ in γ-Ni_{1-x}Fe_xOOH or γ-NiOOH (Fig. 10(b)).¹²⁴

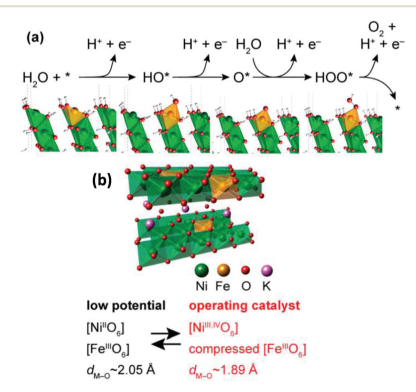


Fig. 10 (a) Representative OER route for NiFe-LDHs and schematic models of the structural transformation of NiFe-LDH into γ -Ni_{1-x}Fe_xOOH through the OER potential, and the corresponding OER path for NiFe LDHs explains the involvement of HO*, O*, OOH* intermediates and Fe-doped γ -NiOOH as active sites. (b) Structure model of Fe-doped γ -NiOOH. Reproduced with permission from ref. 124, Copyright 2015, American Chemical Society.

As mentioned above, NiFe-based materials possess characteristic properties that drive the OER. Most importantly, the exposure of more active sites of NiFe-LDH with enhanced surface area is another approach to increasing the OER performance. Accordingly, Zhang et al. reported the morphological variation from the bulk NiFe-LDH to hollow microsphere (HMS) for better OER activities.84 Bulk LDH possesses the intrinsic characteristics of metal hydroxides, with low electrical conductivity that affects their catalytic properties. Song et al. exfoliated bulk NiFe-LDH to single layer NS by a liquid-phase exfoliation technique for superior OER activity.125 However, the complexity of the liquid phase exfoliation has drawn more attention towards the exfoliation free approach. Han et al. designed a single-layer NiFe-LDH NS by intercalating MoO₄²⁻ ions through the hydrothermal reaction and achieved satisfactory OER performances.¹²⁶ Nevertheless, the poor electrical conductivity of NiFe-LDH and the slow movement of the metal hydroxide layers (Ni(OH)2, Fe(OH)3), drastically affected the mass transfer in the OER. Therefore, NiFe-LDHs are more prone to combining with highly conductive graphene as a support material for the enhancement of the electrical conductivity and the dispersion ability to promote accessibility towards electrochemically active metal centers for catalytic reactions. Following this approach, Long et al. reported the heterostructured NiFe-LDH/rGO hybrid by the electrostatic coupling of the positively charged NiFe-LDH NS with the negatively charged GO layers (Fig. 8(a)).¹¹² As a result, this catalyst displayed outstanding OER activity in alkaline media, with the lowest overpotential of 0.195 V among the noble-metal-free catalysts, and a Tafel slope of 39 mV dec⁻¹, which is close to that of the Ir/C catalyst. These outcomes are due to the homogeneous carbon network of GO/rGO with large surface areas and the uniformly distributed active sites of NiFe-LDH, respectively.

Intercalated graphene has a tremendous role in enhancing the electrocatalysis of NiFe-LDHs. By utilizing the concept of intercalated graphene, Sasaki et al. designed the NiFe/GO superlattice heterostructure via the heteroassembly of NiFe-LDH NS with GO NS by a coprecipitation method using anthraquinone-2-sulfonate (AOS) and hexamethylenetetramine (HMT) as the oxidizing and hydrolyzing agents, respectively, as described in the synthesis section (Fig. 9(a)).⁸³ The morphology of the NiFe/rGO superlattice shows the periodic stacking of NiFe-LDH with GO/rGO, as revealed by the high-resolution TEM/HRTEM image (Fig. 11(a and b)). The graphene, especially for the more conductive rGO layers intercalated into the NiFe-LDHs, is useful for the improvement of electrocatalytic activities of the NiFe-LDHs/rGO superlattice (Fig. 11(c and d)). It displayed a low overpotential of 0.21 V with a Tafel slope of 40 mV per decade, a bit higher than that reported by Long et al., i.e., NiFe-LDH/rGO (0.195 V, 40 mV dec⁻¹).¹¹² Overall, the improved OER performances of NiFe-LDH/rGO

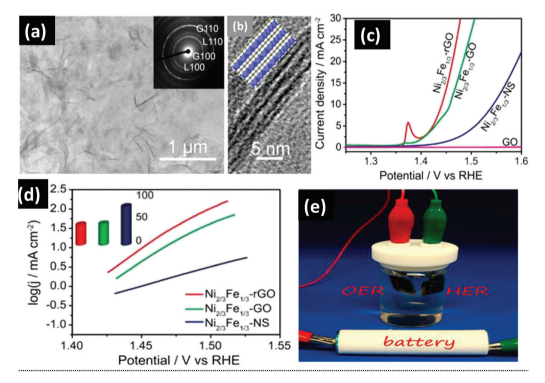


Fig. 11 (a) TEM image of $Ni_{2/3}Fe_{1/3}$ -NS/GO composites. (b) HRTEM image of $Ni_{2/3}Fe_{1/3}$ -NS and GO composites. (c) OER activity of $Ni_{2/3}Fe_{1/3}$ -NS, $Ni_{2/3}Fe_{1/3}$ -GO, and $Ni_{2/3}Fe_{1/3}$ -rGO with IR-compensation. (d) Tafel plot of $Ni_{2/3}Fe_{1/3}$ -NS, $Ni_{2/3}Fe_{1/3}$ -GO, and $Ni_{2/3}Fe_{1/3}$ -rGO. (e) Water-splitting electrolytic cell power-driven by 1.5 VAA battery. Adapted with permission.⁸³ Copyright 2015, American Chemical Society.

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have been attributed to the intercalation of GO/rGO, which restricts the self-aggregation of NiFe-LDH NS and further enhances the exposure of more electrochemical active sites of NiFe-LDH NS. Also, the face-to-face contact between NiFe-LDH and GO exerts stronger interfacial interactions for the electrocatalytic OER process. However, the authors also established an electrolyzer cell fueled by a battery of 1.5 V AA for overall water splitting (Fig. 11(e)). The strategy established by Sasaki's group was further applied to NiMn-LDHs NS for superior electrochemical performance.¹²⁷

More importantly, the incorporation of topological defects by the removal of heteroatoms on the graphene surface is of great importance in accelerating the electrocatalytic water splitting reactions. This defective graphene (named DG) has a greater affinity for the anchoring of transition metal atoms through strong π - π bonding interactions and acts as the active sites for the overall water splitting and oxygen reduction reactions.¹²⁸ Jia et al. designed a heterostructured NiFe-LDH NS/ DG by the electrostatic coupling of NiFe-LDH NS onto defective graphene and used it as a bifunctional electrocatalyst for overall water splitting (Fig. 12(a)).⁹² In particular, the OER catalytic activity of NiFe-LDH NS/DG in alkaline solution was enhanced with a small Tafel slope of 52 mV dec⁻¹ and the overpotential was reduced to 210 mV at the current density I =10 mA cm⁻² (Fig. 12(b-d)). NiFe-LDH-NS/DG also performed well in the HER with a Tafel slope of 110 mV dec⁻¹ and a reduced overpotential of 115 mV at the current density of 20 mA cm⁻² in 1 M KOH (Fig. 12(e and f)). Moreover, NiFe-LDH-NS/DG showed excellent stability towards the OER (Fig. 12(d)) and HER (Fig. 12(g)) in 1 M KOH electrolyte. The advantages of coupling NiFe-LDH-NS/DG for outstanding overall water splitting performances are as follows: (i) DG with

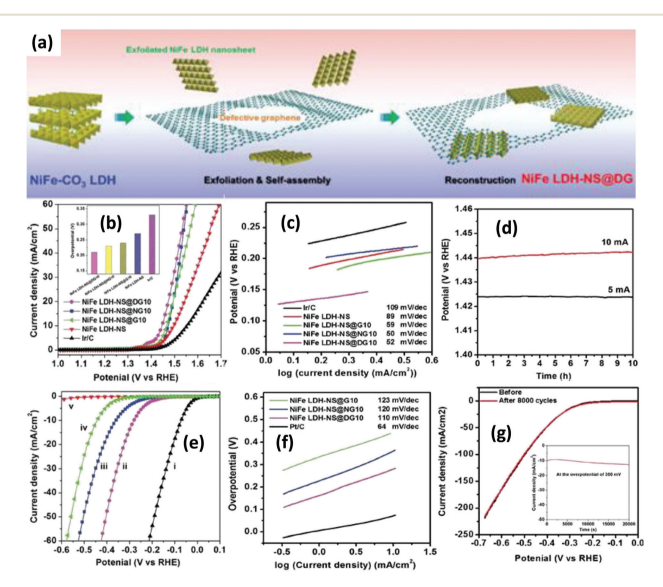


Fig. 12 (a) Illustration of the synthetic route to the NiFe-LDH-NS/DG hybrid. (b) OER LSV curves of the NiFe-LDH-NS/DG series and related materials in 1 M KOH. (c) OER Tafel plots. (d) Stability testing of the NiFe LDH-NS@DG10. (e) HER LSV plots (f) HER Tafel plots. (g) Polarization versus chronoamperometric curves of the NiFe LDH-NS@DG10. Reproduced and modified from ref. 92, Copyright 2017, Wiley VCH.

defective sites provides superior conductivity and high surface area that facilitates the formation of a heterostructure with NiFe-LDH NS for maximum direct interfacial interaction. (ii) The heterostructure formation causes interfacial contractions between Ni/Fe atoms and DG, which speed up the electronic transportation and shorten the diffusion length. (iii) The good dispersion of NiFe LDH NS on DG exposes all active sites of NiFe-LDH for electrocatalytic advantages.

The NiFe-LDH/graphene heterostructures discussed above are promising materials for electrocatalytic water oxidation. Nevertheless, the hydroxides reportedly show inferior HER activity as compared to phosphides and chalcogenide materials. In contrast, the LDH/MoS₂ superlattice has received much attention because the 1T-phase MoS₂ not only behaves as a conductive framework imparting conductivity to the heterostructure, it also shows HER activity similar to commercial Pt/C. Motivated by this, Xiong et al. prepared the MoS₂/ NiFe-LDH superlattice via a solution-phase assembly by restacking the two oppositely charged NS (Fig. 13(a)).¹¹⁰ When MoS₂ and LDH NS were restacked in a layer-by-layer assembly, significant radical charge distribution by electron accumulation at the interfacial area and electron transfer among adjacent monolayers occurred, which synergistically promoted the adsorption of intermediates both in the OER and HER (Fig. 13(b)). The transfer of electrons from the LDH to MoS_2 was 0.45 Å⁻², which revealed the strong electronic coupling between metallic MoS₂ and NiFe-LDH NS as compared to that for LDH/graphene (named LDH/G) and MoS₂/graphene (MoS₂/ G) superlattices. The modulated electronic environment of MoS₂/LDH increased the adsorption between the intermediates and electrocatalyst. Specifically, the localized electron or hole accumulation at the basal edges of 1 T MoS₂ or the LDH NS counterparts resulted in the improved affinity for H⁺ or OH⁻, respectively, which are favorable for efficient OER and HER. This MoS₂/LDH superlattice acts as a bifunctional electrocatalyst, displaying enhanced activity as compared to the other two superlattices (Fig. 13(c and d)). In particular, MoS₂/ LDH exhibited superior HER performance to the MoS₂/graphene composite and unilamellar MoS₂ NS. Most importantly, the rating of MoS₂/LDH as a bifunctional catalyst for the overall water splitting even surpasses the $RuO_2(+)||Pt/C(-)|$ couple as shown in Fig. 13(e). The molecular-level modulation of the heterointerface between the 2D LDH and other unilamellar NS could be of great promise in advanced electrocatalysis.

As mentioned previously, the superlattice structure can effectively modulate the electronic environment of a material. To further optimize the electronic coupling as well as the adsorption energy, the presence of 3D porosity in NiFe-LDH/ rGO could be another strategy for enhancing the catalytic activities of nanomaterials. Porosity increases the active sites on the catalyst surface as well as mass transfer.¹²⁹ Zhan *et al.* reported a porous nanostructure of NiFe-LDH/rGO *via* a facile two-step process involving the reverse micelle solvothermal method and chemical reduction (Fig. 14(a)). The porosity of the NiFe-LDH/rGO nanospheres is due to the aggregation of

the LDH hexagonal plates to produce numerous defective microspheres owing to the reverse micelle effect caused by the role of surfactants as templates.93 Broken GO nanoplates are then attached to the LDH surface, cross-linking it with their oxygen-containing functional groups to generate a rough porous surface as evident from TEM and HRTEM images in Fig. 14(b and c).⁹³ The NiFe-LDH/rGO are highly active electrocatalysts for the OER and ORR, providing excellent OER catalytic activity with a low onset overpotential of 250 mV and a Tafel slope of 91 mV per decade (Fig. 14(d and e)). These enhanced dual catalytic tendencies may have arisen from the surfactant-induced porous microstructure and the sturdy bonding effect of rGO with NiFe-LDH, which are favorable for improved conductivity, more exposure of active sites, and both mass and charge transport of oxygen towards electrocatalytic reactions. Alternatively, the conductive graphene support for the FeNi-LDH generates a 3D framework that is interconnected with the catalytically exposed sites and adequate micro/macroporous channels for fast electron transportation, which facilitate the gas molecule and electrolyte diffusion for the OER. Zhang et al. used a single-step hydrothermal method to introduce NiFe-LDHs into the 3D rGO framework, with conductive CNT (FeNi-GO/rGO LDH) as an additive for further application in the OER.117 The most outstanding OER performance of NiFe-LDH/rGO had a Tafel slope of 39 mV dec⁻¹ and overpotential of 206 mV. In this illustration, FeNi-rGO/LDH demonstrated greater OER performance, possibly due to the nucleation and growth strategy of NiFe-LDH on GO/rGO and further hybridization through the interlayer of NiFe-LDH, which facilitated the disclosure of the active sites in the catalyst, thus increasing the charge transfer kinetics through the rGO layers. By forming NiFe-LDH/rGO, tuning the intrinsic properties of the NiFe-LDH layers triggered the incrementally exposed surface of NiFe hydroxide, resulting in an inimitable layered structure that could effectively encourage the OER performance.

For a better deposition of NiFe-LDH on the graphene surface, the open porous structures of the 3D graphene frameworks having higher conductivity, surface defects, flexibility, and lightweight are desirable factors. As discussed in the growth strategy, Shi *et al.* described a double-step approach to designing a freestanding NiFe-LDH/3DErGO electrode (Fig. 5(c)).⁷⁸ The co-operative result was that NiFe-LDHs and 3D-ErGO substrates formed an exceptional OER catalyst with a low Tafel slope of 39 mV dec⁻¹ and very low overpotential of 259 mV at 10.0 mA cm⁻². The crucial role of Fe in the NiFe LDHs phase was observed with a drastic reduction of the Tafel slope and overpotential with the doping content of Fe (Fig. 14(f-i)).

Recently, Feng *et al.* assembled a strong system of 3D ternary electrodes by the *in situ* growth of vertically aligned $Co_{0.85}Se$ NS on delaminated graphene foil, followed by the growth of NiFe-LDHs by a simple hydrothermal reaction, which displayed excellent performance in the OER with a 250 mA cm⁻² current density at a small potential of 1.51 V.¹³⁰ Heterogeneous atom doping is a significant strategy for enhan-

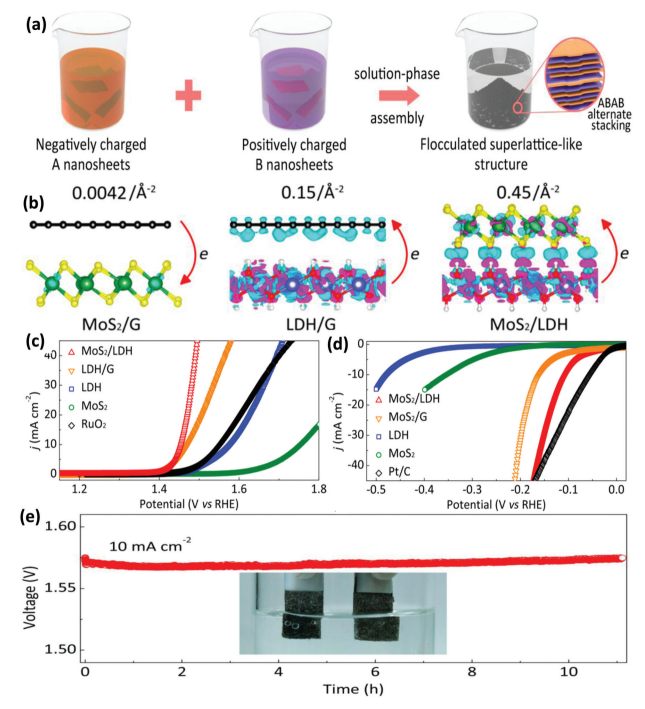


Fig. 13 (a) Schematic of the fabrication of the superlattice by the solution-phase assembly. (b) Bilayer charge distribution by the Bader charge analysis of the MoS_2/G , LDH/G and MoS_2/LDH superlattices. Electrocatalytic performance of MoS_2/LDH and related material for (c) the OER and (d) HER. (e) Evaluation of the stability and overall water splitting activity by the MoS_2/LDH superlattice as the bifunctional electrocatalyst. Adapted with permission.¹¹⁰ Copyright 2019, American Chemical Society.

cing the properties of graphene-related nanomaterials, owing to the positive co-operative effect among heteroatom dopants, carbon atoms and transition metals.^{131,132} Zhang and Wei *et al.* designed N-doped graphene frameworks with spatially confined hybridization with nano NiFe-LDHs (nNiFe-LDH/NGF).¹⁰⁶ In this work, the tremendous catalytic activities in

the OER were proved by the remarkable lowering of the overpotential (\sim 337 mV at 10 mA cm⁻²) and a calculated Tafel slope of 45 mV dec⁻¹ with prolonged durability, which indicate the significance of the work for better electrocatalysis. This work is of importance because of the electron donor properties of N-doping and N-possessing active sites to trigger

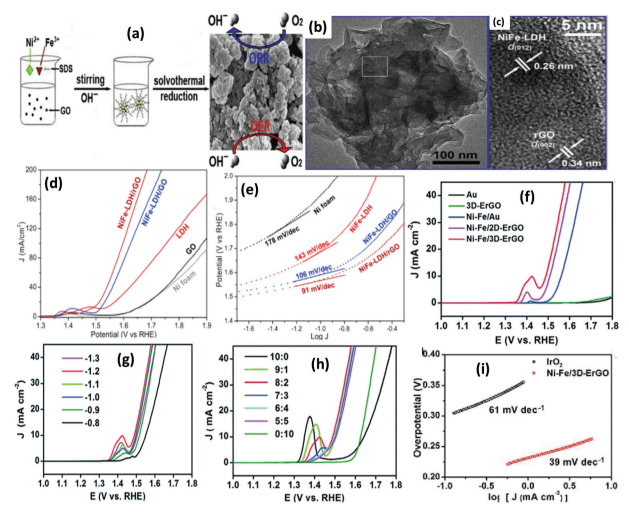


Fig. 14 (a) Synthesis of NiFe-LDH/rGO composites. (b) TEM images of the NiFe-LDH/rGO composites. (c) HRTEM image of the NiFe-LDH/rGO composite. (d) OER activity on Ni foam, GO, LDH, NiFe-LDH/GO, NiFe-LDH/rGO. (e) Tafel plots of Ni foam, GO, LDH, NiFe-LDH/GO and NiFe-LDH/rGO electrodes. Adapted with permission.⁹³ Copyright 2016, Elsevier. (f) LSV curves of Au, 3D-ErGO, NiFe/Au, NiFe/2D-ErGO, and NiFe/3DErGO electrodes. (g) LSV curves of Ni–Fe/3D-ErGO electrodes prepared at the NiFe deposition potential ranging between –0.8 and –1.3 V. (h) LSV plot of the NiFe/3D-ErGO from the variation of Ni/Fe ratios in the precursor solution. (i) Tafel plots of NiFe/3D-ErGO and IrO₂.⁷⁸ Adapted with permission from the Royal Society of Chemistry.

carbon atoms in graphene material for the creation of a large number of defects or active sites to augment the electrocatalytic activities.

The 3D porous structured graphene, with high surface area and porosity and the homogeneous distribution of the porosity in support materials, created a contact surface between the reactant molecules and electrochemically active centers for faster mass transfer. Along with the in-depth investigation into the oxygen reduction reaction (ORR) catalysts, layered structures with pores and interconnectivity would certainly increase the density and contact of extremely sensitive sites.¹³³ Therefore, the design of 3D graphene aerogels is of great importance to the optimization of the mass transfer as well as the exposure of the extremely sensitive active sites of a catalyst. Very recently, Manna *et al.* designed nanostructured NiFe-LDH decorated over the N-doped entangled graphene framework (NEGF), by an in situ solvothermal and freeze-drying method at low temperature and high vacuum pressure.⁷⁶ The freeze-drying method significantly contributes to the formation of a stable entangled graphene network as shown in Fig. 15(a). The synergistic effect of NEGF in the strong coupling with NiFe-LDH facilitated a drastic reduction in the electrocatalytic OER overpotential of NiFe-LDH/NEGF to 290 mV, with a small Tafel slope of 68 mV dec^{-1} in alkaline medium, which is similar to the recent record (Fig. 15(b and c)).⁷⁶ In EC water splitting, the role of graphene in the NiFe-LDH/graphene system is to optimize the mass transfer process by participating as a support material through the 2D-conductive substrate, the open porous structure of the 3D graphene frameworks, the 3D porous entangled structure or free-standing structure, and the 3D defective graphene support, respectively.

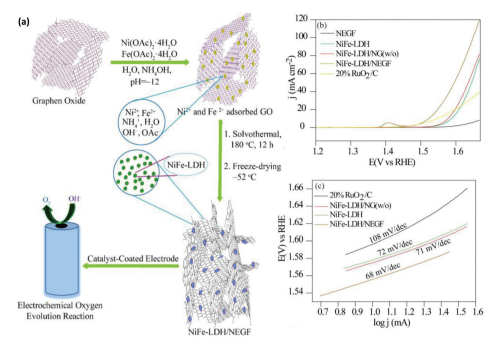


Fig. 15 (a) A pictorial depiction of the synthetic steps involved in NiFe-LDH/NEGF and the corresponding OER activity. (b) Polarization activity of NEGF, NiFe-LDH, NiFe-LDH/NG (w/o), NiFe-LDH/NEGF, and 20%RuO₂/C. (c) Tafel plots of NiFe-LDH/NEGF, NiFe-LDH, NiFe-LDH/NG (w/o), and 20% RuO₂/C.⁷⁶ Adapted with permission from the Royal Society of Chemistry.

Photoelectrochemical (PEC) water splitting

PEC cells are capable of converting direct energy from sunlight into chemical energy. Inside the water splitting reaction, H₂O is oxidized by minority charge carrier holes at the photoanode, releasing electrons and protons into the solution, with the reduction reaction occurring at the cathode, liberating H₂. PEC water splitting into H₂ and O₂ by using solar energy combined with a solar energy collector and water electrolysis in a solitary photoelectrode, which could perform both light absorption and the water splitting reaction. Semiconductors are generally used as photoelectrodes in PEC water splitting reactions. When an n-type semiconductor having a higher chemical potential (calculated by Fermi level alignment) is in contact with the electrolyte, the flow of electrons takes place from the semiconductor to the solution unless the chemical equilibrium is maintained in the solution (Fig. 16(a and b)). In this method, charge transport generates positive depletion or space-charge-region correlating interfacial electric fields that stabilize the difference in potentials among the electrodes and electrolyte solution. However, an unsymmetrical population of carrier charge is established between the VB and CB edge of semiconductors with the irradiation of quantum of energy greater than the bandgap energy of the semiconductor (Fig. 16(c)). This causes the formation of a quasi-Fermi zone between the electrons of the CB ($E_{\rm F}$, n) and the holes of the VB $(E_{\rm F}, p)$. This potential difference across the Fermi levels of the junction, the inbuilt photovoltage (E_{pv}) , is the supreme force

for the water oxidation reaction (Fig. 16(c)). Nevertheless, an external potential bias (E_{app}) is still needed for superior charge separation by varying the inner electric field and mass transport by an external circuit through the transportation of electrons formed in the anodic reaction to the cathode. However, the external potential bias required in PEC is much smaller than that of the EC cell.

The NiFe-LDH/graphene hybrid was employed to facilitate the separation of exciton pairs and thus advance the photocurrent density of PEC electrodes with improved kinetics of the O₂ evolution reactions. The synergistic effect of N-doping and oxygen vacancies can contribute to the superior catalytic activity and stability towards PEC water oxidation.⁶⁰ As discussed in the electrocatalytic OER section, the combination of the N-doping and 3D porous structure could be another weapon for effective charge separations in the NiFe-LDH/graphene nanohybrid. Accordingly, Hou et al. designed 3D hybrid aerogels by mixing N-deficient porous g-C₃N₄ NS (DPCN), N-doped GO/rGO (NRGO), and NiFe-LDH NS via the hydrothermal and freeze-drying methods (Fig. 17(a-d)).⁶⁰ The morphology of DPCN/NRGO/NiFe-LDH revealed the interconnected 3D hierarchical structure and the elemental composition was verified by energy-dispersive X-ray (EDX) spectroscopy. The spatial separation of photogenerated exciton pairs and the unique 3D porous structures with micro-pores favoring mass transfer are the reason this ternary DPCN/NRGO/NiFe-LDH catalyst showed a higher photocurrent density of 162.3 µA $\rm cm^{-2}$ at 1.4 V with unusual photostability after more than 10 h under AM 1.5 G exposure (Fig. 17(e and f)).⁶⁰ Moreover, the charge transfer mechanism revealed that the photogenerated

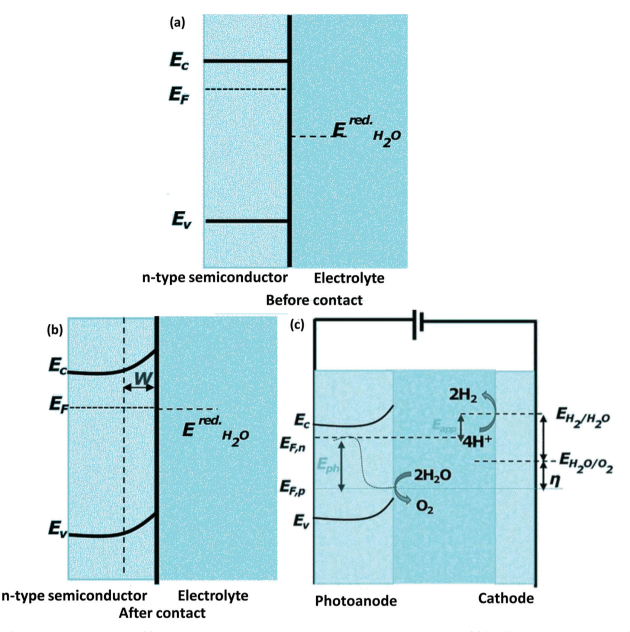


Fig. 16 Schematic representation of (a) the interface of the semiconductor and electrolyte before contact. (b) Equilibration of the interface of the semiconductor–electrolyte after contact. (c) PEC water splitting mechanism under irradiation; $E_c = CB$ edge potential, $E_v = VB$ edge potential, $E_F =$ Fermi edge, $E_{app} =$ applied potential, $E_{pv} =$ photovoltage, W = depletion area, $E^{red}H_2O =$ redox potential of H_2O or the electrolytic solution.

CB electrons of DPCN easily migrated towards the 3D NRGO network. Consequently, the 3D NRGO behaved as an electronconducting transport "highway" that further drifted the electrons toward the current collector. Meanwhile, hole transfer takes place from the VB of DPCN to the NiFe-LDH surface of the NRGO interlayer, which facilitates water oxidation reactions. This process justified the synergistic catalytic effect by constituent semiconductors towards the high-performance PEC water oxidation activity.

Generally, the physicochemical properties of LDHs can be exploited by constructing different hierarchical structures as inspired by nature. The rational design of hierarchical nanostructures such as core-shell, NAs, and nano-pyramids could be an efficient approach to enhancing the PEC water splitting. Hierarchical nanostructures of NiFe-LDH/GO could form 3D channels for rapid electron and hole separations. Han *et al.* designed the BiVO₄/rGO/NiFe hydroxide photoanode that encompasses hierarchical nanostructures by using a facile annelation–photoreduction–electrophoretic deposition method.⁸⁸ The morphological features of this ternary nanocomposite reveal a nanopyramid pattern with the NAs of BiVO₄ wrapped in the interlayer mixing of graphene NS and amorphous NiFe hydroxides, achieving the highest current density of 1.30 mA cm⁻² at 1.23 V. The boost in current

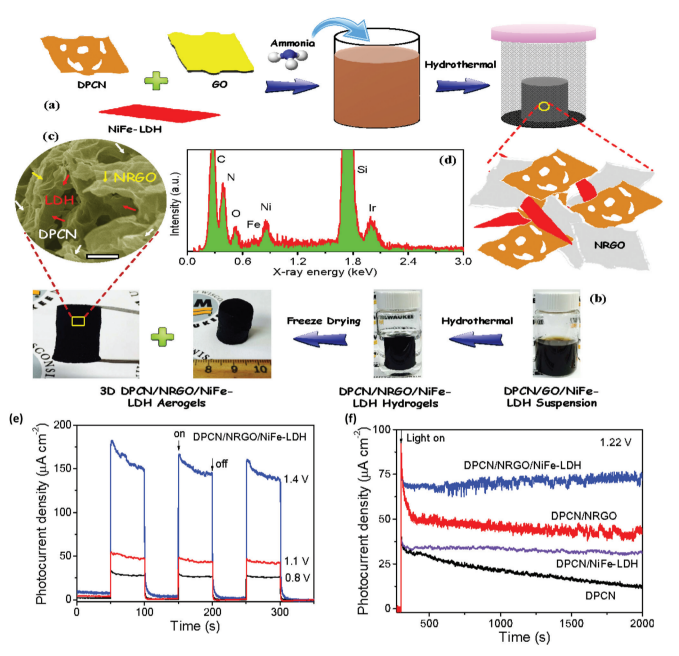


Fig. 17 (a) Schematic representation of the synthetic steps of DPCN/NRGO/NiFe-LDH. (b) Snapshots of the aerogel. (c) FESEM images of DPCN/ NRGO/NiFe-LDH. (d) EDX spectrum of DPCN/NRGO/NiFe-LDH. (e) Photocurrent *vs.* time plot of DPCN/NRGO/NiFe-LDH at different potentials under AM 1.5G irradiation. (f) Stability plots of DPCN, DPCN/NiFe-LDH, DPCN/NRGO, and DPCN/NRGO/NiFe-LDH under AM 1.5G exposure at 1.22 V. Adapted with permission.⁶⁰ Copyright 2016, American Chemical Society.

density can be achieved only due to the synergistic effects among the shuttling of the graphene interlayer as the midlayer in promoting the separation and transfer of photoinduced exciton pairs with the co-catalytic effect of the NiFehydroxide NA as an outer layer, which enhances the stability and PEC activity of the photoanode.

Interestingly, the incorporation of rGO as a solid-state mediator in a p–n heterostructure catalyst plays a great role in the efficient charge separation and the enhancement of the PEC activity of the photoanode. The role of rGO as a mediator

in the NiFe-LDH/GO coupled system provides efficient charge transfer among rGO and NiFe-LDH. In this concept, Sun *et al.* designed a triad BiVO₄/rGO/NiFe-LDH photoanode *via* a two-step facile electrodeposition method as shown in Fig. 18(a).¹³⁴ The morphology of the BiVO₄/rGO/NiFe-LDH photoanode revealed the good contact of the NiFe-LDH NS on the BiVO₄/rGO heterojunction (Fig. 18(b–d)), which facilitated the hole transfer in the PEC water oxidation. More importantly, the triad photoanode exhibited a current density of 1.13 mA cm⁻² at 1.23 V with a broad absorption in the visible zone

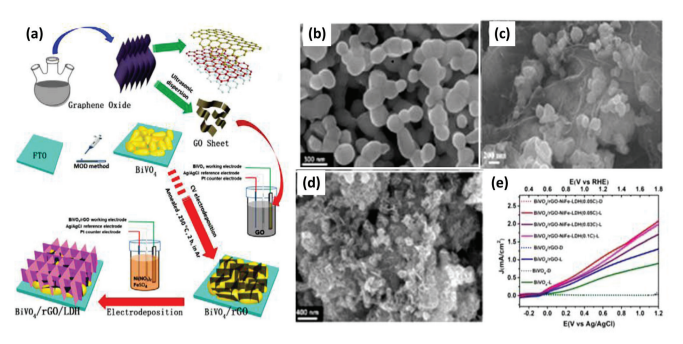


Fig. 18 (a) Schematic representation of the synthetic steps involved in $BiVO_4/rGO/NiFe-LDH$. SEM images of the as-synthesized (b) $BiVO_4$, (c) $BiVO_4/rGO$ and (d) $BiVO_4/rGO/NiFe-LDH$. (e) Photocurrent density of $BiVO_4/rGO/NiFe-LDH$ with the variation in the loading quantity of LDH.¹³⁴ Adapted with permission from the Royal Society of Chemistry.

(Fig. 18(e)). The supreme PEC properties of the triad photoanode were due to the following factors: (i) the creation of the p-n rGO/BiVO₄ heterojunction, (ii) the role of rGO NS as an effective mediator in the shuttling of electrons, and (iii) the role of NiFe-LDH NS as a cocatalyst in speeding up the transfer of holes during the PEC water oxidation.

Following the above major factors, Fig. 19 explains the detailed mechanism of the charge transfer in the $BiVO_4/rGO/$

NiFe-LDH triad photoanode. BiVO₄, a narrow bandgap semiconductor, drastically suffers from bulk recombination. In BiVO₄/rGO, the fast photoinduced electrons of BiVO₄ migrate towards rGO NS as a "mediator with electron shuttling" power that radiates from its delocalized electrons in the sp² bonded carbon network, which then increases the charge carrier lifetime in the BiVO₄/rGO photoanode. When NiFe-LDH as a cocatalyst couples with BiVO₄/rGO then accumulates holes at the

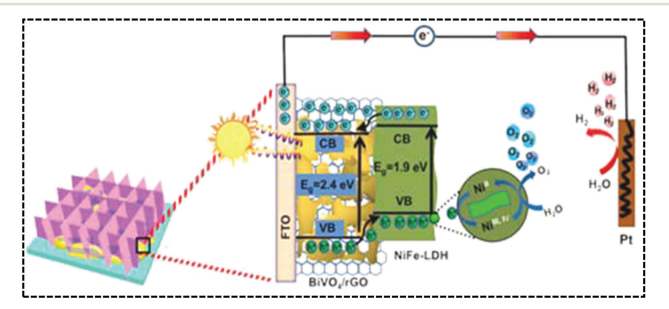


Fig. 19 Illustration of the PEC water oxidation mechanism by the BiVO₄/rGO/NiFe-LDH ternary photoanode.¹³⁴ Adapted with permission from the Royal Society of Chemistry.

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surface of NiFe-LDH, it is known as a hole collector that can facilitate water oxidation reactions. It is actually the holes of BiVO₄ that participate in the redox cycle of the Ni cations of NiFe-LDH to release oxygen. Thus, a low-cost photoanode with high PEC activity and robustness is observed in BiVO₄/rGO/NiFe-LDH as compared to the hierarchal nanopyramid BiVO₄/rGO/NiFe hydroxide photoanode. Based on the PEC water splitting section, the different roles played by graphene in the NiFe-LDH/graphene system for enhancing the PEC activities could include an electron sink, conductive electron highway, and mediator for electron shuttling.

Photocatalytic water splitting

In 1972, Honda and Fujishima performed pioneering work on PEC water splitting using a TiO₂ electrode immersed in an aqueous electrolyte solution and irradiated with UV light source. The photogenerated holes were used to oxidize H₂O to form O₂ and electrons were used to reduce H₂O to form H₂ on a Pt counter electrode via the application of an external potential by a chemical bias or power supply.¹³⁵ This effort has motivated researchers regarding overall water splitting using powder particle photocatalysts for the stoichiometric evolution of H₂ and O₂ as perceived in 1980 by Lehn et al.,¹³⁶ Sato and White et al.,¹³⁷ and Domen et al.¹³⁸ However, the UV active nature of the TiO₂ photocatalyst has its limitations. Researchers have directed their interest towards water splitting using visible light-active semiconductor photocatalysts to harvest a wide portion of visible light since 4% of sunlight is UV light and the rest is visible light. Many groups have reported visible light-active photocatalysts such as Bi₂O₃, WO₃, BiVO₄, CdS, etc. Although these materials have established promising performances through intensive studies, there are some limitations such as short lifetimes, toxicity, instability due to photocorrosion or self-oxidation, and low efficiency towards visible light absorption, which restrict their practical applications.^{5,139,140} Regarding photocatalytic performance, efficiency and stability, the development of more advanced, robust and extremely competent semiconductor photocatalysts for photocatalytic water splitting is still a challenge and more efforts are needed for the advancement of robust photocatalysts.¹⁴¹⁻¹⁴³ Heterogeneous photocatalytic water splitting is a green technology that introduces dispersed powdered photocatalysts and is more beneficial than the PEC technique because of its facile approach, sustainability, cost-effectiveness and external potential.^{33–37,44} Generally, semiconductor photocatalysts used in photocatalytic water splitting depend on several necessities concerning the bandgap energy and electronic properties as follows. (i) The photoactive semiconductor must possess a band structure, inside which the differences in the position of the CB and VB create a bandgap and should harvest the visible light of the solar spectrum; (ii) it should be capable of separating photoinduced e⁻ from reactive h⁺; (iii) a minimum amount of energy should be dissipated during charge transportation and recombination by photoinduced

charge carriers; (iv) there should be chemical stability in aqueous media; (v) it should be kinetically active for electron transportation from the surface of a photocatalyst to the water interface, and (vi) possess appropriate band edge potentials for overall water splitting e.g., the CB of a photocatalyst should be situated at a higher negative potential than the H₂O reduction potential ($H^+/H_2 = 0 V \nu s$. NHE, pH = 0). Similarly, the VB of a photocatalyst must surpass the H₂O oxidation potential to facilitate the oxidation reaction (+1.23 V vs. NHE, pH 0). Consequently, a theoretical semiconductor bandgap energy of $\Delta G = +237$ kJ mol⁻¹ or 1.23 eV is required to force the overall water-splitting (eqn (1.6)-(1.8)). This amount of energy is comparable to a photon energy of $\lambda = 1010$ nm, with bandgap (eV) = $1240/\lambda$ (nm). Hence, theoretically, *ca.* 70% of each solar photon is accessible for water splitting.¹³⁶ All these steps are critical for photocatalytic water splitting reactions. Thus, a robust photocatalyst must possess the appropriate electronic band structure, surface characteristics and bulk properties to contribute to water splitting. The reduction and oxidation reactions of water splitting are as shown in the following equations:

Oxidation
$$H_2O + 2h^+ \to 2H^+ + \frac{1}{2}O_2$$
 (1.6)

Reduction
$$H^+ + 2e^- \rightarrow H_2$$
 (1.7)

Overall water splitting
$$H_2O \rightarrow H_2 + \frac{1}{2}O_2$$
 (1.8)

Half-reactions of water splitting proceed by using sacrificial agents and are mainly suitable for deciding whether or not a specified semiconductor satisfies the thermodynamic and kinetic requirements for the liberation of H2 and O2. Reducing agents such as alcohol (methanol, ethanol), triethylamine (TEA), triethyloxysulphides (TEOS), sulphites and ethylenediaminetetraacetate (EDTA) have been used as sacrificial agents for water reduction, and oxidizing reagents such as persulphate, AgNO₃, Fe³⁺ have been used as sacrificial agents to facilitate water oxidation. However, overall water splitting by using only pure H₂O in the absence of sacrificial reagents fails to give good results. Alternatively, photoinduced electrons at the CB edge of a semiconductor photocatalyst are attracted by oxidizing agents (electron acceptors or electron scavengers) such as AgNO₃, and irreversibly reduce the oxidizing agent as an alternative to H_2O . They augment holes in the VB of a photocatalyst and enhance the liberation of O2 gas. The following equations are involved in photocatalytic H₂ production using methanol as a sacrificial reagent. The product formaldehyde (eqn (1.15)) is then oxidized to methanoic acid (HCOOH) and then to CO_2 and H_2 as described by eqn (1.16) and (1.17).

Semiconductor Photocatalyst $\xrightarrow{h\nu} e^- + h^+$ (1.9)

$$h^+ + H_2O \rightarrow OH + H^+ \tag{1.10}$$

$$2\mathrm{H}^{+} + 2\mathrm{e}^{-} \to \mathrm{H}_{2} \tag{1.11}$$

$$CH_3OH + OH \rightarrow CH_2OH + H_2O$$
 (1.12)

$$^{\bullet}\mathrm{CH}_{2}\mathrm{OH} \rightarrow \mathrm{HCHO} + \mathrm{H}^{+} + \mathrm{e}^{-} \tag{1.13}$$

$$2H_2O + 2e^- \to H_2 + 2OH^- \tag{1.14}$$

$$CH_{3}OH \xrightarrow{h\nu, Catalyst} HCHO + H_{2}$$
(1.15)

$$\text{HCHO} + \text{H}_2\text{O} \xrightarrow{h\nu, \text{ Catalyst}} \text{HCOOH} + \text{H}_2 \qquad (1.16)$$

$$\text{HCOOH} \xrightarrow{n\nu, \text{ Catalyst}} \text{CO}_2 + \text{H}_2 \tag{1.17}$$

Photoinduced excitonic charge recombination is a major obstacle for photocatalytic water splitting reactions. The concept of heterostructure formation is an effective approach to inhibit the charge recombination and the assembly of the NiFe-LDH/graphene-coupled system is a topic of interest. NiFe-LDH with its strong optical response and unique structure has become a hot material for photocatalytic applications. The metal centers in NiFe-LDH are connected by 'oxo-bridged' linkages,^{144–146} which serve as potential sites for charge separation and trigger the optical response of these materials. Also, the presence of the hydroxyl group oxidizes holes in a homogeneous reaction.⁶² Alternatively, graphene is recognized as a substrate material, carrier, mediator and a cocatalyst for photocatalytic water splitting reactions owing to its higher specific surface area and easier carrier charge mobility.³³ We have also reported different NiFe-LDH-based heterostructure materials for various light-driven photocatalytic reactions such as in g-C₃N₄/NiFe-LDH,²⁷ Ag@Ag₃PO₄/g-C₃N₄/NiFe-LDH,⁶² NiFe-LDH/N-rGO/g-C₃N₄,³³ and MoS₂/NiFe-LDH.³⁴ For the first time, in our work, g-C₃N₄/NiFe-LDH has provided deep insight into the heterojunction/nanocomposite effect of g-C₃N₄/LDH for targeted photocatalytic H₂ and O₂ production. In this regard, the coupling of g-C₃N₄ with LDH semiconductors could be a remarkable approach to improving the photocatalytic performance owing to the 2D-2D layered structure, which is of great advantage for utilizing full electronic coupling for effective charge separation. The morphology and chemical composition of LDHs can be delicately tailored to achieve the targeted charge separation efficiency as well as electrical conductivity.²⁷

Researchers and scientists are quite interested in developing robust catalysts with strong redox and effective charge separation ability that could strike the most productive results in water splitting. As discussed in previous sections, heterostructure formation can only meet these entire criteria. NiFe-LDH/graphene in combination with other semiconductors can form heterojunctions in which graphene can act as a mediator for the spatial separation of exciton pairs while preserving higher redox potentials. However, there are no reports of NiFe-LDH/graphene for photocatalytic water splitting except for our breakthrough work, *i.e.*, NiFe-LDH/N-rGO/g-C₃N₄.³³ In this work, the heterostructured NiFe-LDH/N-rGO/g-C₃N₄ was constructed by a facile calcination–electrostatic self-assembly– hydrothermal method as shown in Fig. 7. N-rGO exhibits immense advancement in electrical conductivity linked with the incorporation of the nitrogen heteroatom and also assumes the qualities of graphene such as the outstanding physicochemical properties and large theoretical surface area. In NiFe-LDH/N-rGO/g-C₃N₄, N-rGO acts as a solid-state electronic mediator, integrated between NiFe-LDH and g-C₃N₄ for superior PC reactions. This catalyst displayed outstanding PC H_2 and O_2 evolution rates of 2508 $\mu mol~g^{-1}~2~h^{-1}$ and 1280 μ mol g⁻¹ 2 h⁻¹, respectively (Fig. 20(a and b)), and also displayed exceptional catalytic performances in degrading 95% of Rhodamine B (RhB) dye and 72% of phenol (Fig. 20(c-e)). The role of graphene in the ternary NiFe-LDH/N-rGO/g-C₃N₄ heterostructure for photocatalytic water splitting can be clarified by analyzing the mechanism of charge separation. The coupling of N-doped rGO with catalytic active sites on NiFe-LDH triggered the dynamics of the charge migration phenomenon at the interfacial area of LDH/g-C₃N₄. In the mechanistic approach, NiFe-LDH NS and g-C₃N₄ NS form strong electrostatic bonds with the N-rGO surface, where the CB electrons of LDH recombine with the VB holes of g-C₃N₄. The N-rGO mediator assists in decreasing the charge transportation time, lengthens the lifetime of charge carriers, and endorses the separation of photoinduced carriers charge pairs. In this way, N-rGO near the flat band potential $(E_{\rm fb})$ of NiFe-LDH and g-C₃N₄ can increase the kinetics of water reduction and oxidation potential of the ternary heterostructure. Such attempts indicate the importance of rGO in the charge transport dynamics next to the interfacial area of LDH/g-C₃N₄, as well as the visible-light-harvesting ability of the heterostructure. As realized from the PC water splitting study, graphene mostly plays the role of mediator in NiFe-LDH/graphene-based systems for superior PC performance (Table 1).

Conclusions and outlook

We have summarized an updated strategy for the design of the high-performance NiFe-LDH/graphene nanocomposites for promising energy conversion applications. NiFe-LDH/graphene nanocomposites possess many advantages as follows: (1) they incorporate the advantages and diminish the shortcomings of the solitary materials. (2) The low electrical conductivity of NiFe-LDH is corrected by the support of the conductive graphene matrix. (3) Intercalated graphene acts as a protective layer, effectively restricting the self-aggregation of NiFe-LDH NS and thereby improving the stability. (4) They expose more catalytically active sites due to the good contact between Ni/Fe atoms and doped graphene. (5) The spatially confined growth of nanosized NiFe-LDH in the mesopores of 3D-graphene opens up many coordinate sites for the catalytic process. In the synthesis strategy, the recent advancement in new synthesis protocols for the NiFe-LDH/graphene hybrid with precise structure, morphology and functions are summarized. The in situ growth strategy summarizes the controlled hybridization of the active phases of NiFe-LDH into the conductive graphene substrate, mostly controlling the composition and morphology under different reaction parameters and the effect on

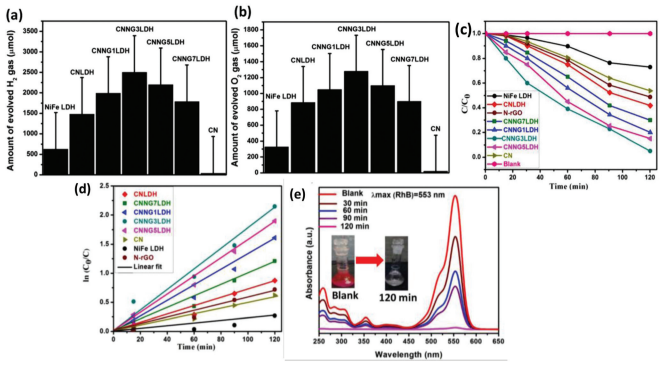


Fig. 20 (a) Plots of the quantity of H₂ evolved (μ mol g⁻¹). (b) Plots of the quantity of O₂ released (μ mol g⁻¹) on NiFe LDH (LDH), g-C₃N₄ (CN), CNLDH (CNNG*x*, *x* = 1, 3, 5 and 7 wt% of GO w.r.t. the quantities of NiFe-LDH (1 g)) and CNNG*x*LDH. Variation of N-rGO on CNLDH, *i.e.*, *x* = 1, 3, 5 and 7 wt% of GO against NiFe LDH in CNNG*x*. Adapted with permission from ref. 33, Copyright 2019 Nature.

the water splitting reactions. The in situ growth also forms strong bonding interactions between NiFe-LDH and graphene, providing the materials with the desired morphologies. Alternatively, the heteroassembly of NiFe-LDH/graphene summarizes the favorable modes of interaction that form a stable and active structure for efficient water splitting; for example, face-to-face contact between NiFe-LDH NS and GO NS through the layer-by-layer approach allows strong interfacial interactions that expose more catalytic active sites for water splitting. It is worth noting that a synergistic effect is realized in the resultant heterostructure and the performances of the NiFe-LDH/graphene nanohybrid, which are fully dependent on the dimensions of graphene materials, the pre-treatment of NiFe-LDH and the reaction conditions. Firstly, the NiFe-LDH/ 2D-graphene composites possess larger specific surface areas, superior electrical conductivity and dimensional orientation, which expose a huge population of active sites of metal centers and boost the charge particles for superior water splitting in EC/PEC/PC. Secondly, the NiFe-LDH/3D-graphene structure exhibits a more axial orientation than the 2D-graphene based nanohybrids and offers a potential substrate for the dispersion of a huge population of catalytically active sites. It also has properties like hierarchical porosity, entangled structure, multidirectional electron mobility and enhanced conductivity, which are responsible for the superior EC/PEC performances.

For an effective energy conversion process, many systematic improvements are needed regarding the fabrication and synthesis of NiFe-LDH/graphene nanocomposites. Accordingly, the research aspects and future prospectives of NiFe-LDH/graphene nanocomposites are as follows:

(i) It is important to directly form 3D freestanding electrodes or grow spatially confined LDH on a conductive 3D graphene matrix due to the creation of more defect sites and enhanced properties that are different from their precursor material, which will facilitate superior electrocatalytic activities with long-term stability; however, there is a long way to go to meet the requirements of the industry. Hence, in-depth studies on the design of free-standing electrodes and coupled catalytic systems using NiFe-LDH/graphene are required for potential applications in electrocatalytic devices.

(ii) The advantages of LDH materials can be partly due to their derivatives such as metal oxides, oxyhydroxides, hydroxides, phosphides, sulfides, bimetal nitrides, and selenides. These LDH derivatives exhibit outstanding EC activities owing to their modulated local electronic environments, enhanced electrical conductivity and improved corrosion resistance properties. However, less work has been conducted on the morphological and topological control in LDH derived mixed metal oxides and nitrides towards water splitting. Moreover, NiFe-LDH and their derivatives are complicated owing to the overlapping of the Ni2+/Ni3+ redox cycle, the OER current density, the onset potential and the Tafel slope. Therefore, there is plenty of room for the optimization of the electronic structure of NiFe-LDH derivatives by computational methods and morphological control by a simple experimental set-up for advanced catalytic properties.

 Table 1
 Recent advances in the water splitting activities of NiFe-LDH/GO-based composite materials

Catalysts	Preparation method	Applications	Electrolyte (mV dec ⁻¹)	Tafel slope	Overpotential	Ref.
NiFe LDH/rGO superlattice	Electrostatic flocculation	Electrocatalytic-OER	1 M KOH	40	$217 \text{ mV} = 10 \text{ mA cm}^{-2}$	83
FeNi-LDH@3DG	Hydrothermal	-OER	0.1 M NaOH	39	$206 \text{ mV} = 10 \text{ mA cm}^{-2}$	117
Ni5Fe1-LDH/RGO	<i>In situ</i> solvothermal	-OER	1 M KOH		$245 \text{ mV} = 10 \text{ mA cm}^{-2}$	66
Ni5Fe1-LDH/rGO	Solvothermal, chemical reduction	-OER	0.1 M KOH	91	$250 \text{ mV} = 10 \text{ mA cm}^{-2}$	93
NiFe-LDH/rGO	Solvothermal	-OER	0.1 M KOH	33	$250 \text{ mV} = 10 \text{ mA cm}^{-2}$	100
NiFe-LDH/NGF	Templated growth, precipitation	-OER	0.1 M KOH	45	337 mV@10 mA cm ⁻²	106
NiFe-LDH/NrGO	Solvothermal	-OER	0.1 M KOH	63	$258 \text{ mV} = 10 \text{ mA cm}^{-2}$	72
NiFe-LDH/3D-ErGO	Electrodeposition	-OER	1 M KOH	39	$259 \text{ mV}(2000 \text{ mM}) \text{ mM} \text{ cm}^{-2}$	78
NiFe-LDHs/NF@3D-rGO	Doctor blåde, freeze-drying, hydrothermal	-OER	1 M KOH	57	$170 \text{ mV} \otimes 20 \text{ mA cm}^{-2}$	75
3D N,S-rGO/WSe ₂ /NiFe-LDH	Hydrothermal, electrostatic self-assembly	-OER	1 M KOH	86	$148 \text{ mV} @ 10 \text{ mA cm}^{-2}$	79
NiFe-LDH/RGO-3 (NFRG-3)	Solid-phase exfoliation	-OER	1 M KOH	49	$250 \text{ mV} \otimes 10 \text{ mA cm}^{-2}$	73
FeNi rGO LDH	Electrostatic attraction	-OER	1 M KOH	39	$195 \text{ mV} = 100 \text{ mA cm}^{-2}$	112
NiFe-LDH/NEGF	In situ solvothermal, freeze-drying	-OER	1 M KOH	68	$290 \text{ mV} = 10 \text{ mA cm}^{-2}$	76
NiFe-LDH-NS@DG10	Electrostatic stacking	SMO-	1 M KOH	52	210 mV (a) 10 mA cm ⁻²	92
Co _{0.85} Se/NiFe-LDH/graphene	In situ hydrothermal	-OER	1 M KOH	57	$270@150 \text{ mA cm}^{-2}$	130
Ni5Fe1-LDH/RGO	<i>In situ</i> solvothermal	Photo-electrocatalysis (PEC, O_2)	1 M NaOH		$0.95 \text{ mA cm}^{-2}, 1.23 \text{ V}$	66
TiO ₂ /rGO/NiFe-LDH	Hydrothermal, electrochemical deposition	PEC, O ₂ evolution	$0.5 \text{ M} \text{ Na}_2 \text{SO}_4$		1.74 mA cm^{-2} , 0.6 V	87
DPCN/NRGO/NiFe-LDH	Hydrothermal	PEC, O ₂ evolution	$0.01 \text{ M} \text{ Na}_2 \text{SO}_4$		$162.3 \ \mu M \ cm^{-2}$, $1.4 \ V$	60
BiVO ₄ /rGO/NiFe-LDH	Electrodeposition	PEC, O_2 evolution	Phosphate buffer		$1.13 \text{ mA cm}^{-2}, 1.23 \text{ V}$	134
NiFe-LDH/N-rGO/g-C ₃ N ₄	Calcination, electrostatic self-assembly, hydrothermal	PC	$0.1 \ \mathrm{M} \ \mathrm{Na}_2 \mathrm{SO}_4$		H_2 evolution, 2508 µmol g ⁻¹ 2 h ⁻¹ O_2 evolution, 1280 µmol g ⁻¹ 2 h ⁻¹	33

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(iii) Understanding the structure and activity in terms of interfacial coupling interactions or interface modulation between NiFe-LDHs and graphene for the formation of a superlattice structure can provide information on NiFe-LDH/ graphene nanocomposites, which is advantageous for catalytic performance and stability. Therefore, in-depth comparisons with other superlattice structures and insight into the catalytic mechanism of NiFe-LDH/graphene are highly recommended. Further advances of NiFe-LDH/graphene toward promising energy storage are needed, and more exploration of NiFe-LDH/ graphene towards photocatalytic and photoelectrochemical water splitting is desired.

(iv) Identification of the active sites in NiFe-LDH/graphene under operando conditions and the mechanism of catalyst action can provide a depth understanding of the structure and activity correlation of the associated catalysts and the design of novel materials with enhanced activities. The in situ characterization of the local electronic structure by using fast-growing powerful instrumental techniques under operando conditions provides fundamental principles for the working catalysts. More specifically, the identification of the most active γ -NiOOH phase from the anhydrous β -NiOOH phase in NiFe-LDH during the OER has long been a debate. Although some studies have made great efforts by combining operando experiments and DFT computational studies to justify the elevated OER performance of the NiFe-LDHs on the atomic level, it is still unclear. Therefore, the systematic identification of the catalytic route and operational active phases of NiFe-LDHs in the NiFe-LDH/graphene model with variations in the atomic ratio, composition and physical structure by using sophisticated online, computational and operando conditions are strongly expected.

Conflicts of interest

There are no conflicts to declare.

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