Perspective on intermetallics towards efficient electrocatalytic water-splitting

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Intermetallic compounds exhibit attractive electronic, physical, and chemical properties, especially in terms of a high density of active sites and enhanced conductivity, making them an ideal class of materials for electrocatalytic applications. Nevertheless, widespread use of intermetallics for such applications is often limited by the complex energy-intensive processes yielding larger particles with decreased surface areas. In this regard, alternative synthetic strategies are now being explored to realize intermetallics with distinct crystal structures, morphology, and chemical composition to achieve high performance and as robust electrode materials. In this perspective, we focus on the recent advances and progress of intermetallics for the reaction of electrochemical water-splitting. We first introduce fundamental principles and the evaluation parameters of water-splitting. Then, we emphasize the various synthetic methodologies adapted for intermetallics and subsequently, discuss their catalytic activities for water-splitting. In particular, importance has been paid to the chemical stability and the structural transformation of the intermetallics as well as their active structure determination under operating water-splitting conditions. Finally, we describe the challenges and future opportunities to develop novel high-performance and stable intermetallic compounds that can hold the key to more green and sustainable economy and rise beyond the horizon of water-splitting application.

1. Introduction

The rising global demand for energy is experiencing major challenges with its exponential growth and nearly entire dependency on fossil fuels. Depletion of these natural resources and the inconvenient impact on climate with its
carbon emissions stipulate alternative strategies in energy supply. In the last few years, tremendous efforts have been dedicated to fulfill the required energy demand by renewable and green technologies and are expected to continue to grow in the years to come. In this regard, hydrogen ($H_2$) is considered as an abundant and clean fuel that can be used as a chemical energy store with a minimal loss of energy. $H_2$ has the highest chemical energy per mass with 143 MJ kg$^{-1}$ (ref. 6) and its energy density is three times higher than that of diesel or gasoline (47 MJ kg$^{-1}$). Once produced, $H_2$ is a clean synthetic fuel, energy supplier for households and the economy, and an important raw material for the chemical industry. A practical and attractive approach to produce inexpensive, reliable, and highly pure $H_2$ is electrochemical water splitting. Depending on the pH value of the electrolyte, water splitting is realized with a polymer electrolyte membrane (PEM) electrolyzer in acidic conditions, neutral water or with alkaline electrolyzer (AEC), and at high temperatures with neutral water with solid oxide electrolysis (SOEL). Generally, these technologies are referred to as water electrolyzer cells (WEC) and considered as the most promising technologies for power-to-(P2G) gas conversion. WECs have their own characteristic merits and shortcomings depending on the process conditions, efficacy, materials, and purity of the product. So far, SOEL has shown the highest efficiency of the WEC technologies but demands challenging material properties due to its high-temperature process of 700–900°C.

Alternatively, PEM technology provides high energy efficiency and fast $H_2$ production rate with high purity, however, impedes their catalyst scope mainly to precious metal and metal oxides resulting in the high cost and limiting their widespread industrial application. In comparison to PEM, an AEC is considered the mellow approach and has widely been used as the leading technology in large-scale industrial applications. Mediating $H_2$ evolution in alkaline media allows replacing noble metals with earth-abundant materials, that either are not stable in acidic conditions or only show poor activities. Therefore, AECs have attracted research to explore various 3d transition metals as alternative catalysts for benchmarking noble-metal IrO$_x$, RuO$_y$, and Pt systems.

Over the years, to bring the WECs to cost parity, numerous classes of cost-effective electrocatalysts such as metal oxides/ (oxy)hydroxide, chalcogenides, pnictides, phosphates, borophosphates, borides, selenides, carbides, alloys, and their heterostructures, have been reported either as cathodes or anodes and a majority of them have shown promising results in comparison to the state-of-the-art catalysts. The reason for superior activity is often described as the presence of a higher number of active sites on the surface, better reaction kinetics as well as the modified electronic properties. In this regard, intermetallic compounds possess complex structures with distinct chemical bonding and have already been found applications in magnetism, superconductivity, shape-memory effects, $H_2$ storage, and recently even for heterogeneous catalysis. Due to covalent bonds, intermetallics are considered as (electro-)chemical stable with high electric conductivity. The high activity is often related to a high number of active sites resulting from the highly ordered structure with voids. These enhanced attractive physical, chemical, and electronic properties with suitable element specificity in intermetallic compounds have attracted considerable interest as ideal electrocatalysts for water splitting.

In this perspective, we focus on the most recent advances in electrocatalytic water-splitting with intermetallic materials (Scheme 1). First of all, we present the fundamental understanding of electrocatalytic water splitting and its evaluation parameters. Next, in order to give the readers a conception of the development methodologies, we discuss the underlying aspects as well as advanced synthetic strategies in detail.

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Scheme 1 Intermetallics for water splitting describing the synthesis strategies to access intermetallic materials with different morphological, structural, chemical, and physical properties with diverse elemental composition. This includes insights into active structure and the reasons for their high efficiency and stability in electrochemical water splitting.
2. Electrochemical water-splitting

For hydrogen production by electrochemical conversion of water to oxygen and hydrogen, the free energy of $\Delta G = 238 \text{ kJ mol}^{-1}$ is needed to mediate the reaction (eqn (1)).

$$\text{H}_2\text{O(}l\text{)} \rightarrow \text{H}_2(g) + \frac{1}{2}\text{O}_2(g)$$

(1)

Although it is seemingly a simple straightforward reaction adjunct to plain electrochemistry, the energy loss entangled to mechanistic intricacy demands strenuous efforts to facilitate water splitting.$^{1,31}$

2.1 Fundamentals

Water splitting is divided into two half-reactions, water reduction at the cathode for the hydrogen evolution reaction (HER) and water oxidation for oxygen evolution reaction (OER) at the anode.$^{1,33}$

Cathode (HER):

$$2\text{H}^+ + 2e^- \rightarrow \text{H}_2 \quad E_{\text{Red}}^0 = 0.00 \text{ V}$$

(2)

Anode (OER):

$$\text{H}_2\text{O} + \frac{1}{2}\text{O}_2 + 2\text{H}^+ + 2e^- \quad E_{\text{Ox}}^0 = 1.23 \text{ V}$$

(3)

OER in particular requires significantly more energy than HER due to comparably more intermediates and reaction steps.$^{14}$ Accounting for the sluggish uphill reaction, OER is still referred to as the bottleneck of overall water splitting, and that is why, in the last few years, immense efforts have been dedicated to seek alternative materials that can effectively reduce the kinetic limitation of OER and enable optimal reaction conditions.$^{32}$ Both reactions take place at the electrode surface (anode and cathode) together with the electrolyte and require consideration of the inner and outer Helmholtz layers.$^{35}$ The performance of these reactions can be measured separately using electrochemical techniques, which can be used to determine the exact values of the applied potentials.$^{36}$ In this regard, the electrochemical HER and OER can be expressed in terms of two redox pairs. In OER, the reduced form $\text{H}_2\text{O}$ and the oxidized form $\text{O}_2$ form the redox couple ($\text{O}_2/\text{H}_2\text{O})$. The HER is composed of the reduced form $\text{H}_2$ and the oxidized form $\text{H}^+$ to form the redox couple ($\text{H}^+/\text{H}_2$). Each of both redox systems have different degrees of reducing and oxidizing power as described in the eqn (2) and (3)$^{3}$ and can be described electrochemically by the redox potential $E$ of the redox system.$^{2}$ The redox potential can be described mathematically by the Nernst equation (see eqn (13)).$^{37}$

$$E = E^0 + \frac{R \cdot T}{z \cdot F} \ln \frac{c_{\text{Ox}}}{c_{\text{Red}}}$$

(4)

The constant $z$ describes the number of electrons occurring in the redox system and $R$ is defined as the general gas constant.$^{38}$ The standard potentials are characteristic of each redox system and cannot be determined experimentally. They form a parameter for the strength of the reducing or oxidizing effect of such a system and thus indicate whether electrons are taken up or released.$^{39,40}$ Only the total potential of a galvanic element can be measured and the potential difference between two redox pairs can be determined to a reference.$^{36}$

2.2 Reaction mechanisms

The HER is kinetically favored and takes place relatively thermodynamically uninhibited with the formation of the single intermediate $\text{M-H}_{\text{ads}}$ which requires ideally a binding energy of 0 V.$^{41,42}$ Its kinetics are defined by two intermediate steps with three possible pathways, the Volmer, and the Heyrovsky or Tafel step with two-electron transfers (see eqn (5)–(7), Fig. 1a).$^{2,43,44}$

Volmer:

$$\text{M} + \text{H}_2\text{O} + e^- \rightarrow \text{M-H}_{\text{ads}} + \text{OH}^-$$

(5)

Heyrovsky:

$$\text{M-H}_{\text{ads}} + \text{H}_2\text{O} + e^- \rightarrow \text{M} + \text{H}_2 + \text{OH}^-$$

(6)

Tafel:

$$\text{M-H}_{\text{ads}} + \text{M-H}_{\text{ads}} \rightarrow 2\text{M} + \text{H}_2$$

(7)

The second step depends on the surface concentration of active sites. On surfaces with low concentrations of active sites, the Heyrovsky reaction is preferred (eqn (6)). In this case, a second water molecule is adsorbed at the same active site followed by electron transfer and desorption of dihydrogen and a hydroxyl ion.$^{46}$ The Tafel reaction is the dominant mechanism on catalyst surfaces with a high concentration of active sites. Chemical desorption of dihydrogen is mediated by two active sites close to each other. The distance between the two active sites should not be more than the van der Waals radius of two adsorbed hydrogen atoms, the closer they are the more likely this reaction path occurs.$^{45}$

The mechanism of the OER is thermodynamically more demanding than for HER with complex kinetics that varies depending on the pH of the electrolyte and the catalyst material (Fig. 1b).$^{46}$ The fourfold is the limiting factor in this reaction, which contains at least three intermediates.$^{46,47}$ There are
several pathways discussed in the literature for varying mechanisms under alkaline and acidic conditions.\cite{40,41,42,43} For the OER under alkaline conditions, the oxide pathway or the electrochemical oxide pathway with two active sites and single-site catalysis have been established (eqn (8)-(10)).\cite{44,45}

\[
\begin{align*}
M + \text{OH}_{\text{aq}}^{-} & \rightarrow M\text{-OH}_{\text{ads}} + e^- \\
M\text{-OH}_{\text{ads}} + \text{OH}_{\text{aq}}^{-} & \rightarrow M\text{-O}_{\text{ads}} + H_2O + e^- \\
M\text{-OH}_{\text{ads}} + M\text{-OH}_{\text{ads}} & \rightarrow M\text{-O}_{\text{ads}} + M + H_2O + e^- \\
M\text{-O}_{\text{ads}} + \text{OH}_{\text{aq}}^{-} & \rightarrow M\text{-OOH}_{\text{ads}} + e^-
\end{align*}
\] (8) (9.1) (9.2) (10)

The last step is often described as proceeding concerted, but can also be divided into two intermediate-steps and proceed via the M–OOH$_{\text{ads}}$ intermediate, which determines the overall reaction rate (eqn (11) and (12)).\cite{40,45}

\[
\begin{align*}
M\text{-OOH}_{\text{ads}} + \text{OH}_{\text{aq}}^{-} & \rightarrow M\text{-OO}_{\text{ads}}^- + H_2O \\
M\text{-OO}_{\text{ads}}^- & \rightarrow M + O_2 + e^-
\end{align*}
\] (11) (12)

The proposed mechanisms for one or two active sites differ only slightly and include the same intermediates as M–OH$_{\text{ads}}$ and M–O$_{\text{ads}}$.\textsuperscript{40,45} The main difference is the final O–O bond formation and release of dioxygen.\textsuperscript{40} The reaction via two catalytically active sites take place through two M–O$_{\text{ads}}$ intermediates with a direct combination of both to form 2M and O$_2$ (eqn (13)).

\[
M\text{-O}_{\text{ads}} + M\text{-O}_{\text{ads}} \rightarrow 2M + O_2
\] (13)

Ideally, the binding of the intermediates requires 1.23 eV each, so that free energy of 4.92 eV is required for all four proton-coupled electron transfer (PCET) steps.\textsuperscript{4} All formed intermediates (M–OH$_{\text{ads}}$, M–O$_{\text{ads}}$, M–OOH$_{\text{ads}}$, M–OO$_{\text{ads}}$) influence the kinetics of the reaction and have a limiting effect on the reaction rate (higher overpotential) if the adsorption or desorption or formation of the respective intermediate proceeds energetically unfavorably.\textsuperscript{52} The route via a single active site with the formation of the M–OO$_{\text{ads}}$ intermediate as discussed above is rarely found in the literature but is taken into account for investigations of kinetic barriers during OER.\textsuperscript{52} Despite these differences, there is mutual consensus about the crucial importance of bond interactions in M–O within these intermediates (M–OH$_{\text{ads}}$, M–O$_{\text{ads}}$, M–OOH$_{\text{ads}}$, M–OO$_{\text{ads}}$). The M–O bond characteristics of the active species M are essential for the overall electrocatalytic activity.\textsuperscript{40}

2.3 Activity evaluation criteria

The activity parameters are the most important criteria to evaluate the activity, efficiency, and stability of a catalyst. The overpotential of an electrochemical process is the additional potential required to catalyze the intended reaction at the thermodynamically determined reversible potential ($\eta^*$) under ideal conditions.\textsuperscript{53} Typically the overpotential $\eta$ at a current density of 10 mA cm$^{-2}$ ($\eta_{10}$) derived from CVs or LSVs is used as a benchmark to compare the activities of electrocatalysts and corresponds to the ~10% solar-to-chemicals efficiency.\textsuperscript{54,55} On the other hand, because of the tremendous growth in the field, researchers have now started to evaluate and compare $\eta_{100}$ and $\eta_{500}$ in addition to $\eta_{10}$ which is highly recommended if a catalyst reaches high current densities. In this way, the evaluation of a catalyst activity would be closer to industrial application standards.

The description of the catalyst activity via the Tafel slope provides a lot of information about the properties of the catalyst material and is recommended to measure under steady-state conditions.\textsuperscript{56} The Tafel slope can provide insights into the dynamics of the catalytic processes taking place on the surface and provide information about kinetics and inhibitions during the water oxidation catalysis.\textsuperscript{52} It indicates how much the
potential has to increase in order to increase the resulting (measured) current $j$ by an order of magnitude. It is an indication of how efficiently and dynamically an electrode or the catalyst applied to it, reacts to an applied potential and generates a catalytic current. This also takes into account any changes in the mechanism with different $\eta$ and can be used to determine which of the PCET reaction in the mechanism is the rate-determining step.$^{5,45,57}$

During the HER, the Volmer step is the RDS when the reaction is defined by the adsorption and discharge of H$^+$ on the catalyst surface, which is indicated by a Tafel slope of $\sim120$ mV dec$^{-1}$ (eqn (5)).$^{46}$ In the case of a low $H_{ads}$ concentration on the surface, the Heyrovsky step is dominant (eqn (6)). A second H$^+$ will be adsorbed at the same active site followed by discharge and desorption of H$_2$ and is indicated by a Tafel slope of $\sim40$ mV dec$^{-1}$. If the $H_{ads}$ concentration on the surface is high, a direct combination of the intermediates is possible to generate H$_2$ (eqn (7)) and resulting in a Tafel slope of $\sim30$ mV dec$^{-1}$. For the OER process, a Tafel slope of 120 mV dec$^{-1}$ indicates that the overall reaction kinetics is dominated by the coordination of the hydroxide step (eqn (8)).$^{57}$ A Tafel slope of 40 mV dec$^{-1}$, indicates the RDS of the electrochemical formation of M-O$_{ads}$ (eqn (9.1)).$^{57,58}$ Alternatively the M-O$_{ads}$ formation can proceed via two metal centers indicated by a Tafel slope of 30 mV dec$^{-1}$ (eqn (9.2)). If the reaction is determined by the formation and desorption of $O_3$, a Tafel slope of 15 mV dec$^{-1}$ is expected (eqn (13)).$^{57}$

On the other hand, theoretical modelling has suggested that a Tafel slope of 120 mV dec$^{-1}$ is not necessarily accompanied by the Volmer step as RDS. Kinetic studies based on microkinetic analysis gave the same Tafel slope for different elementary steps based on intermediate coverage on the surface and mass transport effects. Especially considering newly proposed radical coupling mechanisms rather than a single-site mechanism for OER.$^{22}$ Further, Liu and co-workers suggested based on calculations that the thermodynamic-kinetic model is also highly dependent on factors such as binding environments, temperature, and electrolyte pH that should also be accounted into the design strategy for catalysts.$^{29}$

Electrochemical impedance spectroscopy (EIS) is also a powerful tool to investigate mechanisms in electrochemical reactions, charge transfer processes in materials, and surface properties of electrodes.$^{60}$ It provides very precise results on the electrical conductivity of a material and is a frequently based technique.$^{48}$ From the obtained impedance spectra, conclusions can be drawn about the ability of materials to store electrical energy and transfer electrical charge.$^{84}$ The electrodes are measured with a two or three-electrode set-up where the potential transmits an alternating potential with varying frequency ($\omega$) to the sample. By generating a signal proportional to the generated current. An analyzer then determines the impedance $Z$ of the system from the alternating current flowing through the sample and the alternating voltage generated by a generator.$^{84}$

Similarly, the determination of the electrochemically active surface (ECSA) is an in situ method, which determines the number of active centers responsible for the respective reaction on the surface ($A_{ECSA}$) and thus their size.$^{62-64}$ Other ex situ methods such as BET measurement (BET = Brunauer Emmett Teller) to determine the surface area by adsorbing gas molecules such as nitrogen ($N_2$) can be problematic particularly in the case of porous structures since the accessibility to the interior of the structure depends on the size of the adsorbed molecule.$^{65}$ Surfaces determined with $N_2$ can be smaller than with krypton or water and provide different results than surfaces determined with ECSA.$^{62,66}$ Both analytical methods to evaluate the surface of the catalyst have their own merits and disadvantages and should be considered carefully. While the BET does not necessarily correspond to the ECSA and could result in an unfair comparison between catalysts with different surface densities of active sites, the ECSA analysis suffers from inaccuracies due to the yet non-trivial determination of the roughness factor.$^{53}$ Therefore, it is highly recommended to represent activity plots normalized to both BET and ECSA. In addition to the activity indicators, the long-term stability of the system is also a crucial parameter considering their use for commercial applications. This is typically measured via cyclic voltammetry (CV), galvanostatic chronoamperometry (CA) or potentiostatic chronopotentiometry (CP) electrolysis measurement.

Another valuable insight is the faradaic efficiency (FE), which is defined as the efficiency of electron transfer provided by the external circuit to promote the electrochemical HER or OER reaction. At a constant current density applied for a certain period of time of the experiment, a gas sample is taken with a gas-tight syringe and analyzed with a gas chromatograph (GC) calibrated for H$_2$ or O$_2$. The faradaic efficiency is then calculated from the volume of the generated gas during electrocatalysis, in comparison to the current over time and gives directly correlates the number of electrons needed to generate a mole of gas.$^{67-69}$ The turnover frequency (TOF) is another important descriptor to evaluate the catalytic activity which provides the generated H$_2$ or O$_2$ molecules per second at a single active site. However, the precise determination of TOF remains challenging due to the complexity involved in identifying the total number of such active sites and therefore, only a rough estimation is possible.$^{18}$

By calculations of density functional theory (DFT), potential materials suitable for HER and OER can be identified and sorted according to their catalytic activity based on the adsorption ($\Delta E_H$) and binding energy ($\Delta G$) of the above-discussed intermediates.$^{70,71}$ According to the derived calculations from the DFT, a close relationship between the overpotential and $\Delta G$ of the surface adsorbed intermediates can be expressed. With a few exceptions, DFT provides good predictions about the OER and HER activity of the materials.$^5$ For instance, if $\Delta G_{Hads} \approx 0$ then the material should possess the optimal binding strength for the intermediate and is considered as a good catalyst for HER. But if $\Delta G < 0$, $H_{ads}$ is bound too strongly and desorption of H$_2$ is the RDS shifting kinetics via the Heyrovsky or Tafel mechanism.$^{54}$ A $\Delta G > 0$ indicates a too weakly bounded $H_{ads}$ and therefore, the kinetics is limited by the adsorption of the intermediate, and the reaction proceeds via the Volmer step.$^{53}$ DFT calculations have also been applied to study the relationships of $\Delta G$ in each individual reaction pathway and the activity for both alkaline and acidic media.$^{72}$
3. Introduction to intermetallics

Intermetallic compounds belong to the class of alloys but are distinct by the difference between the subclasses on how the atoms are ordered in the crystalline phase. The bonding in intermetallics involves the combination of partly ionic or covalent interactions instead of weak metallic bonds making them an ordered lattice. Depending on their structure, alloys are initially classified into two different categories: as a solid solution or as an intermetallic compound (Fig. 2). Solid solutions alloys with metal species of similar atomic radius, electronic character, and crystal structure identical to that of the parent metal form a substitutional solid solution with a statistical distribution of the atoms on the lattice sites (Fig. 2a). If the atomic radius of an element is sufficiently small to fit into the lattice spaces of the parent metal, an interstitial solid solution forms (Fig. 2b). In these cases, the composition in the degree of mixing can vary almost at will. Based on a linear interpolation between the properties of the parent materials, the same physical properties can be expected in the resulting alloy. This situation is different when two metals A and B to form an intermetallic compound (Fig. 2c). Intermetallic phases are specified by a clearly defined stoichiometric composition of the elements such as AB, AB₂, A₃B, or even complex mixtures such as A₆B₂₃. Ideally, the metals A and B are not randomly distributed but are arranged at specific positions in the unit cell and differ for A and B. This often results in physical properties that are different from the physical properties of the composite elements with a well-ordered crystal structure disparate from those of the parent elements. Hence, a semiconductor can be derived from two metallic conductors in an intermetallic phase, or a magnetic intermetallic phase can be formed from non-magnetic composing materials.

Intermetallic systems have the advantage that a heterogeneous catalytic process such as the adsorption of reactants on the catalyst surface or the activation of reactants mediating the reaction to the desired product can be assigned to the suitable incorporated species which can cooperate via so-called interfaces in the composite material. When it comes to composite materials, intermetallics have a particularly large number of interfaces between the different types of atoms in their structure. A particularly pronounced synergism between activation of the reactants on the surface by one metal and realization of the reaction by fine-tuning of the redox potential with the help of the other can be achieved. The synergetic effect of the two metallic components improves the properties compared to the individual homometallic components.

Recently, it has been shown that most of the compounds containing anions/nonmetals are unstable under water splitting conditions (especially in OER) and depending on the testing period, they transform either partially or completely to their corresponding mostly layered oxide/oxyhydroxide structures. Such surface/bulk transformed materials from non-oxidic materials have proven to be more active compared to the bare oxide materials due to either high ECSA leading to higher accessible active sites or better electronic conductivity. As the intermetallic compounds are element specific with complex crystal structures and can be formed by both active and conducting elements, they are naturally considered good candidates for electrocatalytic water splitting.

4. Chemical synthetic strategies of intermetallics

Most of the so far known 6000 binary intermetallics are mainly derived from solid-state techniques since they have been primarily investigated for their physical and structural
properties.\textsuperscript{74,76,78} Intermetallic compounds often show increased stability, selectivity, and activity for a variety of catalytic processes due to their complex structure and bonding characteristics.\textsuperscript{73,75,79} and therefore, over the years, many new advanced synthetic approaches to control their particle size, shape, morphology, and surface area have been developed to gain wider adoption.\textsuperscript{10,77} In the following, several important strategies for the synthesis of intermetallics, starting from single crystals to nanostructures, towards catalytic HER and OER have been discussed in detail.

4.1 High-temperature solid-state methods

A historical approach to the synthesis of intermetallics is the thermal annealing method in a protective atmosphere or evacuated conditions at relatively high temperatures to produce thermodynamically stable products in large single crystals from \( \mu \mbox{m up to \( \mu \mbox{m scale}.\)\textsuperscript{80} Though little control remains of the reaction pathway during synthesis (temperature, pressure), and often exploratory efforts are required to attain the desired phase-pure and functional compounds.\textsuperscript{80,81} Giving access to a broad variety of metallic alloy and intermetallic phases, this method is often used to study materials for their physical, chemical, and mechanical properties.\textsuperscript{82–86} The benign technique to develop and predesign intermetallic precursors for the catalytic application has given rise to a large number of water-splitting electrocatalysts.\textsuperscript{87–90}

For instance, Liasia and co-workers synthesized Ni–Mo\textsuperscript{91} and Ni–Mo–Al\textsuperscript{92} based intermetallic phases for HER of various compositions by mixing the elemental powders in stoichiometric amounts and melted under an inert atmosphere. While mixing Ni with Mo gave phase pure Ni\(_2\)Mo and Ni\(_3\)Mo, the Ni–Mo–Al phase was a mixture of Ni\(_2\)Al\(_3\) or Ni\(_2\)Al\(_3\)/NiAl\(_3\) with NiAl\(_2\)Mo\(_2\).\textsuperscript{91,92} Similarly, a phase pure NiAl\(_3\) was prepared in vacuum at temperatures between 600–1250 \textdegree\textsubscript{C} by the group of Liu using commercially available Ni carbonyl and gas atomized Al powder as precursors. The resulting material of a highly porous morphology showed excellent performances in HER catalysis.\textsuperscript{93,94} Meanwhile, Ni\(_3\)Al prepared by Han et al. in vacuum at a temperature of 1000 \textdegree\textsubscript{C} using the induction melting technique showed good activity during electrocatalytic OER.\textsuperscript{95}

Along this direction, mixing elemental Al and Co metal powders followed by heating in the Ar atmosphere resulted in the formation of an intermetallic nanoporous Al\(_x\)Co\(_2\) framework that after additional sulfurization acted as a highly efficient pre catalyst for HER.\textsuperscript{96}

Motivated by the previous results, we recently synthesized polycrystalline Mn\(_3\)Ga\(_4\) by annealing a stoichiometric mixture of Mn and Ga in an evacuated quartz ampule to 900 \textdegree\textsubscript{C} for four days to ensure homogeneity of the mixture, then cooled down and annealed at 380 \textdegree\textsubscript{C} for further ten days.\textsuperscript{97} Identical conditions were also chosen to prepare Fe\(_3\)Ge\(_5\) from Fe powder and Ge chips where the stoichiometric amount of metals were sealed at 1000 \textdegree\textsubscript{C} for two days before annealing again at 650 \textdegree\textsubscript{C} for seven days.\textsuperscript{98} Both intermetallic phases were examined for catalytic OER in alkaline conditions that showed significant activity and stability.\textsuperscript{97,98} In addition, through solid–gas reaction from a mixture of copper and nickel powders, a highly conductive antiperovskite-based hybrid Cu\(_{1–x}\)NNi\(_{1–y}\) was synthesized and after Fe\textsuperscript{99} treatment, the formation of a p-Cu\(_{1–x}\)NNi\(_{1–y}\)/FeNiCu with a core–shell structure was observed. Such rational design of the catalyst displayed high conductivity and porosity resulting in a remarkable activity towards OER (Fig. 3a).\textsuperscript{100}

The solid-state techniques have been further extended for the preparation of intermetallic phases containing five or more metal elements with high entropic order. Jin et al. were successfully synthesized Al\(_x\)NiCoIr\(_x\) and Al\(_{0.9}\)NiIr\(_x\) by melting using an induction-melting furnace under Ar protection.\textsuperscript{100} Subsequently, the prepared phases were melted again in a quartz tube and injected onto a spinning Cu roller to prepare the ribbons that were then chemically etched in a 0.5 M NaOH solution to prepare nanoporous high-entropy alloys (np-HEAs) for the HER catalysis.\textsuperscript{100} Very similarly, Ding et al. designed a quinary FeCoNiCrNb\(_{0.5}\) from their corresponding elemental powders. The eutectic high entropy alloy (EHHEA) with a porous nanostructure and high corrosion resistance was successfully applied as a catalyst for OER.\textsuperscript{101}

Apart from using traditional induction furnaces and overcome their limitations, arc-melting became a viable method to rapidly (re)melting and refine structures as well as remove high and low-density inclusions during melting.\textsuperscript{102} It is a quite simple and straightforward technique and has widely been applied to attain various noble, i.e. PtDy\textsuperscript{103}, Al\(_2\)Pt\textsuperscript{104} and non-noble metal-based, i.e. Ti\(_2\)Ni\textsuperscript{97}, TiCo\(_2\textsuperscript{105},\) Ni\(_3\)Ta\textsuperscript{106} intermetallic compounds for HER and OER. Along this line, lamellar nanostructured Ni-Co-Al was prepared by arc melting the homogenized raw materials at 1200 \textdegree\textsubscript{C} under Ar atmosphere that showed convincing results for alkaline HER.\textsuperscript{111} In a similar approach, the noble metal-based Hf\(_{5}\)B\(_{2}\)Ir\(_{3}\) phase was obtained via a two-step process in which the metals were first arc melted in Ar at about 1200 \textdegree\textsubscript{C} for several weeks and then applied through spark plasma sintering (SPS) to retrieve the electrode material for OER catalysis.\textsuperscript{112} Moreover, in a unique strategy, Jian and co-workers synthesized nanoporous hybrid Cu\(_{12–x}\)Co\(_{1-x}\)Al\(_{4y}\) (\(x = 0 \mbox{ or } 3, y = 0 \mbox{ or } 1\)) by the group of Liu using commercially available Ni carbonyl and gas-atomized Al powder as precursors. The resulting material of a highly porous morphology showed excellent performances in HER catalysis.\textsuperscript{113}

4.2 Chemical reduction

A relatively mild approach at ambient pressure and low temperature is the chemical reduction of metallic precursors in adequate solvents. Varying the reducing agents from the low reducing power of ethanol and linoleic acid to strong with NaBH\(_4\) and N\(_2\)H\(_4\) gives control over unwanted reactions where noble metals have dominated the reduction process leading to core–shell structures. With the additional use of surfactants and the specific choice of solvents, various noble and non-noble nanoparticles (NPs), e.g. FeIr\(_3\), NiPd, CoPd\(_2\), CuRh, NiPt, etc.,
can be prepared, with varying size and shape from hexagonal or star-shaped to spherical and even small network-like structures. Compared to solid-state, this approach allows overcoming the kinetics energy barrier in the disorder-to-order transition at lower temperatures (<350 °C) allowing structure transition into a well-defined nanoparticulate system with proposed composition.

To improve the catalytic activity of catalysts for water splitting, the active sites must have an optimal balance between adsorption and dissociation of intermediates, high electronic conductivity, cost, and stability. Therefore, the chemical reduction is an efficient technique to modulate the electronic structure, reduce or inhibit noble metal content, modify morphology and surface on the atomic level.

Taking advantage of this approach, nano particular PdBi and PdPb on Pt were prepared that showed the potential of nanoscale surface engineering by modifying rather inactive materials into highly efficient HER catalysts (Fig. 4). Both systems were attained at relatively mild conditions below 200 °C giving highly ordered NPs with improved intrinsic activity. Concurrently, IrNi and IrPt nanoframes were synthesized by reducing corresponding metal salt precursors and etching the framework. The as-attained IrNi systems encompassed a high ECSA and improved intrinsic activity compared to bare Ir. Not only was IrNi more active than bare Ir towards OER, but it also showed improved long-term stability mediating the challenging reaction in acidic media.
At moderately high temperatures, we prepared the intermetallic FeSn2 phase by reducing a stirred mixture of iron(III) and tin(II) salt in a solution of polyethylene glycol and ethylene glycol under a rapid and constant flow of N2. The growth of intermetallic FeSn2 nanocrystals transpired by the interspersing of Fe into the interlayers of metallic Sn through a Kirkendall process (Fig. 3b). An analogous one-pot reduction was also used to prepare isotropic CoSn2 nanocrystals at 170 °C. Both materials were tested for water splitting catalysis with very high activity for both OER and HER.

The hot injection is another viable reduction technique to synthesize monodisperse NPs, via rapid nucleation, followed by controlled growth of nuclei to form larger particles, and is well developed for the preparation of quantum dots. By varying stabilizing ligands, precursors, reaction time, and temperature, various phases of different sizes and morphologies can be achieved. In this respect, for the synthesis of popcorn-shaped GaPt3 NPs, Lim et al. used Ga(C5H7O2)3 precursor in oleylamine (OLA) and preheated to 200 °C under Ar atmosphere. To this solution, PtI2 was injected, and the mixture was heated up to 280 °C for 30 min followed by an instant cooling in a cold water bath. The intermetallic GaPt3 phase was examined for HER in all pH ranges (alkaline, acidic, and neutral). Alternatively, Lim et al. used the hot-injection method to prepare GaPd2 nanomaterials with different morphology such as polyhedrons, NPs, and nanowires just by using different solvents that exhibited both high catalytic HER activity and stability and were far superior to a commercial Pd catalyst. Furthermore, the same group synthesized GePt3 and Ge3Pt and concluded that by tuning the composition of the solvents and reagents as well as the reaction temperature, distinct phases of intermetallic compounds active for HER can be realized. In the meantime, Schaan and co-workers reported a lower-temperature route to colloidal metal silicide NPs such as Pd3Si, Ni2Si, and Cu3Si that were efficient for HER in acidic media.

### 4.3 Hydro(solvo)thermal route

Hydrothermal or solvothermal synthesis is generally defined as a chemical reaction occurring at aqueous or non-aqueous solvents at elevated temperatures above the boiling point of the reaction medium and above atmospheric pressure. This synthetic technique is rather straightforward compared to the high-temperature solid-state and requires mostly moderate temperatures. By controlling the suitable selection of reaction conditions (precursor composition, pH, solvent, heating rate, temperature, pressure, concentration, etc.), pure phases with distinct sizes and morphology can be obtained. Currently, this method has been regarded as one of the versatile and relatively simple approaches to access numerous amorphous and crystalline structures.

Although a variety of intermetallic compounds have been synthesized under such conditions and applied for manifold applications, their utilization as electrocatalysts for water splitting is rather limited. Recently, intermetallic NiMo with a surface containing SiO was obtained in reducing hydrothermally grown NiMoO4 in the H2 atmosphere. This porous NiMo composite was tested for HER with performances close to Pt. In another approach, Jin et al., constructed intermetallic MoNi4 networks for OER, HER, and overall water splitting by treating nickel foam (NF) with ammonium molybdate via a hydrothermal reaction and then post-treated in H2 atmosphere at temperatures between 300–600 °C. Similarly, IrRu@Te intermetallic particles were attained when a mixture of Ir and RuCl3 was mixed together with Na2TeO3 in water and heated for 18 h in an autoclave at 180 °C. Without the addition of Na2TeO3, the unsupported IrRu phase was extracted but showed less activity when tested for OER under acidic and neutral conditions. Furthermore, PdCu3 was prepared solvothermally by mixing Pd(acac)2, Cu(acac)2 and cetyltrimethylammonium bromide (CTAB) together with OLA in a Teflon-lined autoclave at 180 °C for 24 h. The resulting Pd–Cu phase was successfully tested for HER with Pt-like activity.

### 4.4 Wet impregnation

Wet impregnation is a well-known and effective technique for the preparation of heterogeneous catalysts, to significantly influence the physical and morphological properties by altering the microstructure. With suitable precursors, NPs are impregnated into a porous established electrode framework (support) to either enhance the electronic or ionic conductivity of the electrode or to enhance the catalytic activity, if both are not sufficient. Carbon-supported structurally ordered IrV3 NPs with a significantly reduced proportion of Ir and optimal distribution within the structure with a maximized number of neighboring Ir/V bimetallic sites were synthesized from a solution of IrCl3 and VCl3 mixed carbon black. To obtain the intermetallic phase, the well-dispersed suspension was dried and heated at 800 and 1000 °C under H2/Ar (5%) for 3 h. The ordered structure was able to efficiently promote the water dissociation and enhance the kinetics for alkaline HER.
prepared in the same fashion but at a much lower temperature.\textsuperscript{136} The group of Li used N-doped graphite nanotubes as a highly active substrate for their intermetallic Ni–Mo nanocatalysts, (Fig. 3d). First, the Ni–N-doped graphite nanotubes were prepared by pyrolysis followed by acid etching, and then the N-doped graphite tubes with residue Ni (Ni-NGTs) were impregnated in an aqueous ammonium molybdate solution. Under reductive conditions at 600 °C, NGT supported Ni–Mo nanocatalyst was obtained with superior activity and stability in HER catalysis.\textsuperscript{137} Interestingly, recent work of Wu and coworkers revealed that highly efficient HER can also be accomplished by using MXene-supported Pt\textsubscript{5}Ti (Pt\textsubscript{5}Ti\textsubscript{5}C\textsubscript{2}T\textsubscript{2}) through wet impregnation.\textsuperscript{139}

4.5 Single source precursor-derived synthesis

The single-source precursor (SSP) approach is a versatile technique to explore bimetallic or even polymetallic materials. The pre-coordination in metalorganic precursors allows defined assembling of atoms on the molecular level and even altering the chemical behavior giving access to materials that are not approachable in traditional ways.\textsuperscript{140} SSPs enable a relatively good control over the composition, morphology, structural and electronic properties of desired materials. A large number of described metalorganic SSPs provide wide-range access to numerous materials and owing to their low decomposition temperature, various crystalline and amorphous structures have been reported.\textsuperscript{1}

For intermetallic phases, Vela and co-workers compared the SSPs with the conventional precursor approach and they found that the SSP approach is more reliable as different reduction potentials of separate metal ions can be a limiting factor and lead to phase segregation or other unwanted side products.\textsuperscript{21} Further, the SSP techniques provide phase tuning by the change of ligand coordination environment and temperature that is generally lower than solid-state synthesis.\textsuperscript{141,142} Even though SSP is a very convenient strategy to synthesize phase pure intermetallic compounds for water splitting, less attention has been given considering the amount of complexities involved in designing a suitable molecular precursor.

Lately, DiSalvo’s group synthesized an ammonium nickel molybdate with the formula \(\text{NH}_4\text{HNNi}_2(\text{OH})_6[\text{MoO}_4]_2\) and reduced the SSP to attain a Mo\textsubscript{5}Ni\textsubscript{7} and a Ni\textsubscript{0.92}Mo\textsubscript{0.08} phase. Thus yielded intermetallics exhibited promising results for alkaline HER.\textsuperscript{143} Learning lessons from our previous works,\textsuperscript{19,36,38,144–146} we reported a premediated synthesis of xanthene-based bis(germylene)-Ni complex and utilized it as a low-temperature precursor to form monodisperse ultra-small nickel germainide (NiGe) nanocrystals (Fig. 3e). Strikingly, the NiGe exhibited substantially low overpotentials for OER surpassing the state-of-the-art Ni\textsubscript{5}, Fe, Co, and benchmarked NiFe, and noble-metal-based catalysts.\textsuperscript{189} Furthermore, we were also able to access the intermetallic FeAs phase from a novel dinuclear arsenide iron complex \(\text{L}_9\text{FeAs}_2\text{Fe}^9\) (\(\text{L}_9 = \text{CH}(\text{C}^\text{Bu}N\text{Dipp})_9\)) as SSP through a hot-injection method at moderate temperature. FeAs yielded in small spherical particles of about 10 nm in diameter with high surface area and was effective to catalyze OER.\textsuperscript{190} It is important to note that in recent years, SSP-derived materials have attained immense attention for the preparation of novel intermetallic compounds in nano form that otherwise are usually prepared via high-temperature routes and are relatively unexplored for electrocatalytic water-splitting.

4.6 Electrodeposition

A very common strategy to transform molecular precursors into electrocatalysts is the electrophoretic decomposition of these precursors at a certain applied potential to form a coating on a substrate.\textsuperscript{1} Though this strategy often lacks a complete understanding of the correlation between the electrochemical parameters and the resulting microstructure, it is a facile route to access amorphous and nanocrystalline structured films.\textsuperscript{150} Usually, intermetallic coatings protect against corrosion or improve mechanical properties and are of industrial importance.\textsuperscript{151,152}

Pure intermetallic coatings fabricated by electrodeposition without using templates or post-heating techniques remain challenging and only a few examples exist.\textsuperscript{153} But the simplicity of this method and low cost in comparison to other deposition methods yielding in highly active species makes this approach attractive and admirable for application in electrocatalysis.\textsuperscript{154} Intermetallic Fe\textsubscript{5}Zn\textsubscript{9}–FeZn\textsubscript{13} composite as reported by the group of Das or Ni–Zn coatings composed of NiZn, NiZn\textsubscript{3}, Ni\textsubscript{2}Zn\textsubscript{11}, and NiZn\textsubscript{7,31} phases synthesized by Jovic \textit{et al.} proved to be efficient for water electrolysis. The use of simple precursors such as sulfates or chlorides in an aqueous solution to achieve adherent and evenly distributed film at low temperatures makes this approach benign and feasible.\textsuperscript{155,156} On the other end, an approach described by Ballesteros \textit{et al.} resulted in phase pure CuZn\textsubscript{3} films on a Ni foil. Using an aqueous solution of copper and zinc sulfate glycine and a conventional three-electrode set-up, only the size of the spherical crystallites covering the electrode changed depending on the applied potential and the films tested for HER catalysis with moderate activity.\textsuperscript{154} Alternatively, Jovic \textit{et al.} synthesized phase pure NiSn films of varying morphology with varying the current density which showed remarkable activity for OER when tested in alkaline solution.\textsuperscript{157}

4.7 Physical vapour deposition

Physical vapor deposition (PVD) is a widely known and applied technique for film preparation on various substrates based on evaporation or sputtering principles.\textsuperscript{158} This very energy-demanding technology allows precise regulation not only on morphology and layer thickness during film synthesis but also allows material composition far from the thermodynamic equilibrium while forming the targeted species permitting synthesis of ‘new’ materials.\textsuperscript{156,159}

Utilizing this approach, Al–Ni electrodes were fabricated by PVD of aluminum onto a nickel substrate yielding in a coating ~20 μm thick layers on the surface.\textsuperscript{160} The films were then annealed at 610 °C followed by selective aluminum leaching in a strongly alkaline electrolyte. XRD analysis revealed the
formation of a mixture of the intermetallic Al$_3$Ni$_2$ and Al$_3$Ni phases in the films since the Al$_3$Ni$_2$ phase grows faster than the Al$_3$Ni phase. It was proposed that longer heat treatment will lead to solely an Al$_3$Ni$_2$ phase. The prepared electrodes were investigated for activity in alkaline water electrolysis with convincing results but showed severe instability of the materials.\textsuperscript{160}

### 4.8 Other methods

Another convenient route to synthesize intermetallics is the melt-spinning technique. This method has shown great potential in nanostructuring to affect the physical and morphological properties of a material. Especially, for amorphous metallic compounds and due to their capacity of producing nanograins, melt-spinning has been utilized since the early 1980s.\textsuperscript{161} Several intermetallic phases such as FeCoNiAlTi HE162 for HER and Cu/Al$_3$Cu$_4$Ni@Cu$_4$Ni\textsuperscript{163} and NiIrRuAl NPNWs\textsuperscript{164} for overall water splitting have been fabricated via this facile approach to fine-tune their properties for efficient catalysis (Fig. 5).

One of the interesting approaches is high-intensity ultrasound treatment which has recently been used to synthesize the intermetallic AlNi phase. Here, commercially available Al–Ni alloy powder (50 wt% Ni) was dispersed in a water/ethanol mixture and sonicated for 1 h. The acoustic cavitation during the treatment is responsible for the initiation of redox processes on the particle surface leading to changes in its composition. Depending on the water: ethanol composition the different intermetallic Al$_3$Ni$_2$ and Al$_3$Ni phases could be obtained and were successfully investigated for HER catalysis.\textsuperscript{165,166}

Other ingenious methods such as laser ablation or plasma process synthesis have also been utilized to produce PtCo/CoO\textsubscript{x} nanocomposites and FeNi\textsubscript{3} for efficient OER. Lately, the ball milling technique was used to attain Ni$_{10}$Mo$_{40}$Ti$_{10}$ and Ni$_{10}$Mo$_{50}$\textsuperscript{168} for HER and NiFe\textsuperscript{169} for efficient OER electrocatalysis. Based on the stoichiometry mixing the Fe and Ni at different molar ratios, FeNi$_3$ nitrogen-doped carbon (FeNi$_3$/NC) was obtained using a plasma–liquid reaction (Fig. 3c). Plasma engineering improves the electrical properties and surface area and has proven to be a reliable electrocatalyst for OER and ORR.\textsuperscript{181}

Apart from the synthesis, it is also important to thoroughly characterize the intermetallic compounds to understand their geometric and electronic properties so that they can readily be used for electrocatalysis. Usually, powder X-ray and single-crystal X-ray diffraction methods are used to determine their crystallinity, phase identification, and structural analysis with complementary theoretical calculations. Similarly, the surface structural characterization of (amorphous or crystalline) intermetallic compounds are performed by various techniques such as scanning electron microscopy, transmission electron microscopy, selected area electron diffraction, high angle annular dark-field scanning transmission electron microscopy, electron energy-loss spectroscopy, X-ray photoelectron spectroscopy. Besides, the coordination structure of intermetallics is revealed by Raman and X-ray absorption spectroscopy.\textsuperscript{170}

### 5. Intermetallics for hydrogen evolution

Intermetallic phases have attracted many research groups to investigate their unique properties for water-splitting capability. Especially, intermetallic combinations of transition metals, as composite catalysts for HER, have been the subject of manifold experimental and theoretical studies.\textsuperscript{171} The pronounced synergism arises from a combination of transition metals with empty or half-filled d-orbitals with metals having more filled orbitals. The latter transition metals have internally paired d-electrons not available for bonding in the pure metal but in combination, giving exceptional charge-transfer capabilities to the formed intermetallic phase. This leads to exceptional electrocatalytic behavior in their intermetallic bi- or a multi-elemental compound that often surpasses precious metals as described by Brewer–Engel valence-bond theory for bonding in metals.\textsuperscript{172–174}

According to the concept, various intermetallic combinations of transition metals, such as Hf$_2$Fe, Hf$_2$Co, PtMo, Pt–Mo, ZrPt, Nb–Pd systems, PdTa, and TiPt have been used as electrode material for HER in alkaline media and compared with the conventional transition metal-based materials. In contrast to the conventional cathodes, the intermetallics have shown significant electrocatalytic enhancement with Pt$_2$Mo and Hf$_2$Fe as the most efficient HER catalysts.\textsuperscript{175,176}

#### 5.1 Noble metal-based intermetallics

In electrocatalysis, especially, Pt-based intermetallics have shown high catalytic activity and often outperform non-noble-based alloys.\textsuperscript{177} Though in water-splitting, many transition metal-based electrodes have demonstrated high electrocatalytic performance in HER, Pt is still considered the highest active catalyst in alkaline and acidic electrolytes.\textsuperscript{3} With optimal structural properties to adsorb H$^+$ more efficiently, it shows the lowest overpotential and is the most active material in
Enhanced the HER in comparison to the ordered species. 

Formation of Ni/Pt MoPt3 electrode was activated by polishing the surface. Lined a prominent role of Mo in the catalytic process once the uniformly spherical NPs with sizes of 10 sputtering, giving an electrode of addition, a Pt-based silicide was prepared by magnetron-sputtering, giving an electrode of addition, a Pt-based silicide was prepared by magnetron.

Another platinum-based intermetallic, Pt3Ti NPs on a two-dimensional transition metal carbide (Pt/TiC2T2) at different temperatures was investigated recently by Li et al. for HER in 0.1 M HClO4 as the electrolyte. The Pt/Ti NPs prepared at 550 °C (Pt/TiC2T2–550) showed the highest activity in alkaline HER when testing from cooperative effects between the nanowires and stability for 24 h, superior to commercial Pd, was attributed to the increased surface area and the synergetic electron transfer from Pd to Pt facilitating the electrocatalytic HER. 114 115

Although Pt and Ru-based materials have been considered the best working electrocatalysts for HER, very recently, Ir has also garnered significant attention towards HER. 190,191 As Ir can provide better stability at a high anodic potential than Ru in acidic conditions, it is considered as one of the very few catalysts driving efficient OER and HER in a broad range of pH. 191 Making use of the bifunctionality with an oxophilic metal to promote the water dissociation and production of hydrogen intermediates, a structurally ordered intermetallic IrV was synthesized by Chen and co-workers to facilitate the HER. Vanadium being a highly oxophilic metal with the strongest metal–OH bond among the transition metals formed a structurally ordered phase with Ir (IrV) resulting in superior electrocatalytic behavior for alkaline HER. The achieved η–10 was only 9.0 mV even under prolonged conditions and revealed a Tafel slope of 24.1 mV dec–1 and clearly outmatching Pt/C and Ir/C references. 137

5.2 Non-noble metal-based intermetallics

Completely replacing noble metals as catalyst materials remains a challenge in HER, especially since pure Mo, Fe, Co, or Ni bind H+ too weakly or too strong and therefore, are more or less inactive in HER. 192 But when these metals were alloyed, a positive effect on the HER activity was observed with improved H+ adsorption characteristics and reduced overpotentials. 193 Demonstrating exceptional physical and chemical properties, non-noble intermetallics have attracted many research groups to profoundly study this new class of materials for application in HER. 194

In this respect, Rosalbinao et al. revealed that the electrocatalytic properties of Fe could be increased by forming Jana et al. synthesized PdCu3 by chemical reduction in OLA 180 °C and the addition of CTAB to control the reduction rates of Pd and Cu, giving spherical NPs that were highly active towards electrochemical HER. In acidic media (0.5 M H2SO4) high activity of PdCu3 was correlated to the dealloying of Cu from the structure resulting in the formation of active Pd sites with a low coordination number to facilitate the HER. Moreover, in a shape-controlled approach, gallium–palladium (GaPd3) nanomaterials were fabricated to drive the electrocatalytic HER. By changing the surfactant during synthesis, GaPd3 polyhedrons, NPs, and nanowires were achieved which were then examined for HER in 0.5 M H2SO4. The GaPd3 NPs exhibited the highest activity with η–10 of 24 mV and a Tafel slope of 57.2 mV dec–1 followed by the polyhedrons (η–10 of 33 mV) and the nanowires (η–10 50 mV). The enhanced activity of the nanowires and stability for 24 h, superior to commercial Pd, was attributed to the increased surface area and the synergetic effect of Pd and Ga within the material. Lately, the group of Wu decorated intermetallic Pd3Pb nanoparticles with a submonolayer of Pt to enhance acidic HER catalysis. They concluded that the very high activity resulted from the intermetallic substrate that stabilized the atomic structure of the active Pt layer as well as stabilized the electronic structure for effective electron transfer from Pd3Pb to Pt facilitating the electrocatalytic HER. 114

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intermetallic phases of Fe with rare earth metals. By the appropriate combination of 3d-orbitals of Fe with 5d-orbitals of Ce or La, according to the Brewer–Engel valence-bond theory, the HER performance of the final material was significantly enhanced to industrial standard. In their series, they found that the Fe–MM (MM = mixed metal) alloy, primary (Fe) crystals surrounded by a Fe27R2 phase (R content is Ce 5.5, Pr 1.5, Nd 1.5, Sm 0.5, Eu 0.2, Gd 0.3, Tb 0.5 and Dy 0.5 in at%) noted as Fe90MM10 and Fe90Ce10 to be most active for the alkaline HER.99 Furthermore, the positive effect of Ce in the structure was also observed in Ni–Zn-coatings containing the intermetallic NiZn13, Ni2Zn11, and NiZn7.3 phases. The composite material embedded with CeO2 NPs increasing HER activity than the one without the inclusion of CeO2.100

In order to surpass the performance of Raney nickel electrodes, Ti32Ni and TiNi were synthesized which showed considerably good performances in HER with small activation energies to mediate the reaction and extend higher stability than the Raney nickel reference.101–103 Intense studies on the Ti–Ni system revealed a decreasing trend for HER activity in the order TiNi3 > TiNi4 > TiNi5 > TiNi > Ti > TiNi > TiNi0.7, in which only TiNi3 exhibited the highest activity.104 The Raney nickel system with Ni3Al as analogous to TiAl with a similar face-centered cubic unit cell as well as structural order has also been investigated as the catalyst material for HER.105 Early reported phase pure Ni3Al and NiAl3 with minor impurities of Ni2Al3 were tested in 1 M KOH for HER activity and demonstrated good activity and stability. The porosity of the electrode material could be increased with Al content and the addition of Mo in the alloyed material could even further increase the HER activity.106 Comparable observations were made when the intermetallic phases of TiAl, FeAl, and NiAl were additionally alloyed with 2 at% of transition metals M = Ti, V, Cr, Mn, Fe, Co, Ni, and Cu. Additional alloying in general influenced activity, but, the more electron-rich the alloying metal became, the stronger the influence was on the overpotential and the HER activity.107 Similarly prepared Ni3Al by Wu et al. and Dong et al. through elemental powder reaction revealed enhanced porosity of the electrode materials and activity for alkaline HER. They observed that the intermetallic phase showed increased corrosion resistance and acquired a higher surface area due to the porous structure, as well as lower charge transfer resistance enhancing the HER compared to the pure Ni phase.108,109 Besides, Ni-based intermetallic phases combined with rare earth elements such as La or Sm (Mm = mixed metal) also displayed promotional effects on HER activity. Tamura’s group studied LaNi5– and MmNi5-type alloys as electrode materials for HER and found that it shows Pt/Pd-like behavior and activity.110–112 Recently, a Ni–Co–Al lamellar nanostructure was attained by arc melting technique and tested for HER in a 1 M KOH electrolyte. Additional aging of Ni–Co–Al in an oven and etching in alkaline media, the material was dealloyed resulting in a higher ECSA resulting in a higher HER activity with η10 of 178 mV when compared to a Ni foil standard.113 In a very different approach, Sun et al. synthesized Al–Cu2Ni@Cu2Ni core/shell nanocrystals by a melting-spinning method for highly efficient HER in alkaline media. The hybrid material showed η10 of 139 mV and a Tafel slope of 110 mV dec−1 mediating the HER over a prolonged time of 8 h with a minimal shift of ~14 mV. The high activity was accounted to the Ni incorporation that leads to a bimodal nanoporous architecture that simultaneously facilitates electrolyte access and electron transport as well as adapts the binding energy for H+ in nearby Cu atoms.114

Mo-based intermetallics are considered as a suitable substitute for electrode materials in HER since their activity often proved to be much higher with more favorable kinetics than pure Mo or Ni-based electrodes.115–117 McKone et al. directly compared the HER activity of Ni and Ni–Mo nanopowders revealing enhanced per-surface-atom activity compared to the bare Ni.118 Meanwhile, the group Jaramillo investigated intermetallic phases of NiMo, NiMoCo, CoMo, and NiMoFe for HER in acidic conditions and found a much higher activity than for Pt deposited on rotating disc electrodes.119 MoNi3 supported by MoO2 cuboids grown on NF via hydrothermal synthesis and annealing in the H2 atmosphere have shown remarkable activity towards HER, favoring the largely reduced Volmer step. With η10 of 15 mV and a Tafel slope of 30 mV dec−1, this catalyst is considered as the best Pt-free HER catalyst in alkaline media.120 A study on ordered Mo2Ni9, disordered Ni0.92Mo0.08, and pure Ni powders for HER activity was also conducted that revealed the highest activity for the disordered species. Due to the high surface area, the disordered Ni0.92Mo0.08 favored the catalytic process, but when mass-specific activity was normalized to the surface area (determined by BET) rather than interfacial capacitance, the ordered Mo2Ni9 showed more intrinsically activity.121 A positive influence of Ti on the electrochemical performance in intermetallic NiMo was observed by Panek and co-workers. Ni50Mo50Ti10 demonstrated high intrinsic activity and much lower overpotentials for HER than the identically prepared Ni50Mo50.122 Enhancement of HER activity in Ni–Mo electrocatalysts was also achieved by including carbon nanostructures into the catalytic process. By caging Ni5Mo NPs into N-doped graphite tubes, a significant amplification of HER activity was observed with an η10 as low as 65 mV a Tafel slope of 67 mV dec−1 to mediate acidic HER for 15 h.123 Additionally, the catalytic HER activity was improved by covering the surface of a porous NiMo network with N-doped graphene also strengthened the chemical stability of the system.124

Intermetallic stannides are known to possess unique chemical bonding with high electrical conductivity.88 The earlier work on nickel stannides, Ni3Sn and Ni3Sn2, fabricated via solid-state proved to be more active than their homometalllic counterparts.88,89 Likewise, inspired by the previous works, Krstajic and his group electrodeposited Ni–Zn coatings and applied for alkaline HER and acquired promising results. The electrodes contained a mixture of different intermetallic phases (Ni, Ni3Sn, Ni1.2Sn0.8, Ni0.6Sn0.4) with varying chemical composition, phase composition, and morphology, and the effect of morphology on the activity was the most pronounced.137

Most recently, Suen and co-workers systematically synthesized a series of intermetallics, MCo2 (M = Ti, Zr, Hf, and Sc) using a rapid arc-melting method and studied their effects on
HER. In this series, TiCo$_4$ displayed a promising activity with an $\eta_{10}$ of 70 mV, a Tafel slope of 33 mV dec$^{-1}$, and stability of 12 h and, this activity was comparable to a Pt/C standard.\(^{109}\)

Efforts to stabilize and fine-tune the electronic structure by chemical synergism and structural site-isolation have been made to fabricate highly active non-noble catalysts. Based on the concept, FeCoNiAlTi, a high-entropy intermetallic (HEI) possessing the unusual periodically ordered structure was synthesized.\(^{102}\) The partially dealloyed HEI by acid etching showed high alkaline HER activity with an $\eta_{10}$ of 88.2 mV, a Tafel slope of 40.1 mV dec$^{-1}$, and stability of 40 h (Fig. 6a–c).\(^{162}\)

Here, the lowering of the overpotential was ascribed to the chemical complexity and unique atomic configurations that deliver a strong synergistic function to alter the electronic structure by optimizing the required energy barrier for hydrogen evolution.

Apart from conventional intermetallic compounds, sulfur-based intermetallic Al$_{25}$Co$_7$S$_{64}$ was also prepared by dealloying Al$_{90}$Co$_{10}$ in 6 M KOH followed by solid-vapor sulfurization. The Al–CoS$_2$ showed $\eta_{10}$ of 70 mV and a Tafel slope of 153.5 mV dec$^{-1}$ and was found to be stable for 12 h.\(^{36}\) An up-to-date table on the activity parameters of intermetallics has been provided in Table 1.

Although Pt and Pt-based alloys remain as the most active materials for HER with $\eta_{10}$ between 10–20 mV and Tafel slopes ranging from 25–30 mV dec$^{-1}$,\(^{210}\) recently, several non-noble metal catalysts have attracted tremendous attention. Especially, Mo-based catalysts such as CoMoS$_2$ with $\eta_{10}$ = 56 mV and a Tafel slope = 32 mV dec$^{-1}$ or Co–MoC with $\eta_{10}$ = 46 mV and a Tafel slope = 46 mV dec$^{-1}$ running stable for over 500 h, are considered potential candidates to replace Pt in electrochemical HER.\(^{211,212}\) In comparison, judging from the HER activities, the intermetallic-based catalysts have added advantages (see Section 8) and are on par with the other benchmarking systems, thus, are the most suitable candidates for HER catalysis.\(^{213}\)

6. Intermetallics for oxygen evolution reaction

6.1 Noble metal-based intermetallics

Traditionally, RuO$_x$ and IrO$_x$ are still considered as the best working electrode materials for OER but their low abundance and high cost have limited their broad industrial application.\(^{214,215}\) However, the extraordinary stability especially in acidic media still makes these elements the first choice as anodes. Ruthenium catalysts usually are slightly more active and iridium-based catalysts are considered more stable for OER. Taking this into account, intermetallic cata
tals based on Ru and Ir have been investigated both in acidic and neutral OER. Xu et al. reported IrRu@Te accessed via a solvothermal approach that exhibited low $\eta_{10}$ of 220 (Tafel slope = 35 mV dec$^{-1}$), 245, and 309 mV in 0.5 M H$_2$SO$_4$, 0.05 M H$_2$SO$_4$, and neutral electrolytes maintaining continuous OER catalysis in strong acid (0.5 M H$_2$SO$_4$) for 20 h at 10 mA cm$^{-2}$ with a minimal decline in activity.\(^{132}\)

Fig. 6 Electrocatalytic performance for HER and OER of selected catalysts: (a) HER LSV curves and (b) Tafel plots at a scan rate of 5 mV s$^{-1}$ with iR loss correction of FeCoNiAlTI HEI and references in 1 M KOH. (c) Long-term stability measurement of the FeCoNiAlTI HEI at a current density of 20 and 100 mA cm$^{-2}$ without iR loss correction. Figures (a–c) are reprinted with permission from ref. 162, copyright 2020, Wiley-VCH (d) OER LSV curves of Cu$_{1-y}$NNi$_y$/FeNiCu and references obtained from the steady-state measurements. (e) CP measurements of p-Cu$_{1-y}$NNi$_y$/FeNiCu and commercial IrO$_2$ at 10 mA cm$^{-2}$. Figures (d–f) are reprinted with permission from ref. 99, copyright 2020, Nature Publishing Group.
Besides Ru and Ir, nanostructured Pt or prevailing intermetallic phases such as MoPt₃–HfPd₃ composite materials are proposed to provide substantially advanced electrocatalytic properties for OER.²²⁰ To develop corrosion-resistant catalysts for OER, Pt-based catalysts are the optimal alternative to realize highly active and long-term stable electrocatalysis. PtCo NPs embedded into CoOₓ matrices showed remarkable resistance against aggregation and dissolution in alkaline media.²²¹ The

Table 1  Comparison of reported HER overpotentials of selected intermetallic catalysts at a given current densities in alkaline, acidic and neutral media

| Catalyst (substrate) | Electrolyte | Current density (mA cm⁻²) | Overpotential (mV) | Tafel slope (mV dec⁻¹) | Reference |
|----------------------|-------------|---------------------------|--------------------|------------------------|-----------|
| Al₆Cu₅Ni@Cu₄Ni (direct) | 0.1 M KOH | −10 | 139 | 110 | 163 |
| CoSn₃ (NF) | 1 M KOH | −10 | 103 | — | 221 |
| CoSn₃ (FTO) | 1 M KOH | −10 | 196 | 78 | 221 |
| np-AlNiCoIrMo HEA (GC) | 0.5 M H₂SO₄ | −10 | 18.5 | 33.2 | 100 |
| NiIrRuAl-3/1 (GC) | 0.1 M HClO₄ | −10 | 14 | 23 | 164 |
| D-PtNi/C (GC) | 1 M KOH | −10 | 40 | 55 | 138 |
| FeCoNiAtTI HEI (direct) | 1 M KOH | −10 | 88.2 | 40.1 | 162 |
| Fe₉₀Y₁₀ (direct) | 1 M KOH | −250 | 470 | 125 | 89 |
| Fe₉₀MnMn₄ (direct) | 1 M KOH | −250 | 330 | 85 | 89 |
| Ni-Zn (steel) | 1 M NaOH | 2.47 | 0 | 119 | 156 |
| AlNiHT30 + L (Ni) | 1 M KOH | −100 | 123 | 76 | 160 |
| Ni-al (direct) | 6 M KOH | 90.1 | 242 | 219 | 93 |
| NiAl₁₂ @ 25 °C (Cu) | 1 M KOH | −250 | 280 | 200 | 92 |
| NiAl₁₂ @ 25 °C (Cu) | 1 M KOH | −250 | 253 | 157 | 92 |
| NiAl₁₂ @ 25 °C (Cu) | 1 M KOH | −136 | 99 | 92 |
| NiAl₂M₀₂₅ @ 25 °C (Cu) | 1 M KOH | −250 | 57 | 134 | 92 |
| NiAl₂M₀₂₅ @ 25 °C (Cu) | 1 M KOH | −250 | 60 | 99 | 92 |
| Ni-Co-Al (direct) | 1 M KOH | −10 | 178 | 158 | 111 |
| Ni-LaNi₅ (Cu) | 1 M NaOH | 250 | 330 | 101 | 222 |
| Ni-MnNi₃₂Co₂₈Al₆₈ (Cu) | 1 M NaOH | −250 | 351 | 113 | 222 |
| Mo-Ni₂ (Ti) | 1 M KOH | −3.7 (BET) | 200 | — | 143 |
| Ni₀₀₂M₀₂₀ (Ti) | 1 M KOH | −1 (BET) | 200 | — | 143 |
| Mo₅Ni (Ni) | 1 M KOH | −10 | 15 | 30 | 208 |
| Porous MoNi₄ networks (Ni) | 1 M KOH | −10 | 28 | 36 | 131 |
| Co₂Mo (direct) | 1 M KOH | −10 | 12 | 40 | 113 |
| TiCo₂ (direct) | 1 M KOH | −10 | 70 | 33 | 109 |
| HfCo₂ (direct) | 1 M KOH | −10 | 87 | 47 | 109 |
| SeCo₂ (direct) | 1 M KOH | −10 | 87 | 36 | 109 |
| GePt₃ (FTO) | 0.5 M H₂SO₄ | −10 | 53 | 37 | 124 |
| PdCu₃ (GC) | 0.5 M H₂SO₄ | 10 | 50 | 34 | 129 |
| PtSi (Ti) | 0.5 M H₂SO₄ | 40 | 78 | 30.5 | 186 |
| Pt-Dy (direct) | 8 M KOH (@25 °C) | −100 | 409 | 147 | 107 |
| Pt-Dy (direct) | 8 M KOH (@85 °C) | −100 | 139 | 141 | 107 |
| GaPt₃ (GC) | 0.5 M H₂SO₄ | −10 | 27 | 43.3 | 122 |
| GaPt₃ (GC) | 1 M KOH | −10 | 48 | 63.1 | 122 |
| GaPt₃ (GC) | 10 | 103 | 85.3 | 122 |
| PbD₂ (GC) | 0.5 M HClO₄ | −10 | 78 | 63 | 119 |
| GaPdex nanowires (GC) | 0.5 M H₂SO₄ | −10 | 50 | 57.2 | 123 |
| Pd₃Si (Ti) | 0.5 M H₂SO₄ | 192 | 192 | 131 | 125 |
| Ni₂Si (Ti) | 0.5 M H₂SO₄ | 243 | 243 | 66 | 125 |
| Ir½V (GC) | 1 M KOH | −10 | 9.0 | 24.1 | 137 |
| np-CoAl (direct) | 0.5 M H₂SO₄ | −10 | 239 | 153.50 | 96 |
| Pt/C (carbon fiber) | 0.1 M HClO₄ | −10 | 32.7 | 32.3 | 139 |
| AL-Pt/Pd/Pb (GC) | 0.5 M H₂SO₄ | −10 | 14 | 18 | 114 |
| NiMo(a) (GC) | 1 M H₂SO₄ | −10 | 39 | — | 207 |
| NiMo(a) (GC) | 1 M KOH | −10 | 30 | — | 207 |
| Porous NiMo (GC) | 0.5 M H₂SO₄ | −10 | 22 | 37 | 130 |
| NiMo-HG (GC) | 0.5 M H₂SO₄ | −10 | 30 | 41 | 130 |
| NiMo (1 mg cm⁻²) (Ti) | 2 M KOH | −20 | 70 | — | 206 |
| NiMo (3 mg cm⁻²) (Ti) | 0.5 M H₂SO₄ | −20 | 80 | — | 206 |
| NiMo-NCTs (GC) | 0.5 M H₂SO₄ | −10 | 67 | 67 | 104 |
| Ni₉₀Mo₇₀Ti₉₀ (373 K) (direct) | 5 M KOH | −100 | 215 | 249 | 168 |
| Ni₉₀Mo₇₀ (373 K) (direct) | 5 M KOH | −100 | 245 | 240 | 168 |

* FTO = fluorine-doped tin oxide, GC = glassy carbon, NF = nickel foam.
observed $\eta_{10}$ of 380 mV (Tafel slope = 71.2 mV dec$^{-1}$) for OER was far lower compared to Co-oxide and Pt/C with appreciated long-term stability. In a novel approach to assemble intermetallics, Lee and co-workers uncovered rhomboic dodecahedral IrNi and PtNi nanoframes. Within them, IrNi exhibited surprisingly high activity towards OER catalysis with $\eta_{10} = 325.8$ mV a Tafel slope of 48.6 mV dec$^{-1}$ and was structurally stable for 5000 cycles outperforming the Ir/C reference catalyst with a low Tafel slope of 48.6 mV dec$^{-1}$. The enhancement in the activity of IrNi was ascribed to the unique structural and chemical composition of the material. Recently, Antonyshyn et al. synthesized the intermetallic Al$_2$Pt phase for efficient OER giving rise to a moderate $\eta_{10}$ of 450 mV that was better than the Pt reference. In addition to the activity, it showed a stable performance after activation during the first 100 h for more than 450 h.

### 6.2 Non-noble metal-based intermetallics

In the last few years, numerous oxide-based materials have been investigated to address the concerns of the activity and stability of electrocatalysts during OER. Nevertheless, the lack of good electrical conductivity to enable an efficient charge transfer process constrains their widespread use. In this respect, transition metal-based intermetallics have been regarded as the most suitable electrocatalysts for OER as they comprise both desired chemical and physical properties. As intermetallics can also be cost-effectively prepared on a large scale, they have gained enormous attention in the field of water-splitting catalysts. The group of Schuhmann recently investigated several intermetallics such as NiAs ($\eta_{10} = 360$ mV, Tafel slope = 58.7 mV dec$^{-1}$) and Ni$_5$Si ($\eta_{10} = 410$ mV, Tafel slope = 70.9 mV dec$^{-1}$) for noble metal-free OER in alkaline electrolytes. They observed enhanced intrinsic OER activity in several metalloids resulting from unique electronic and structural properties in certain composite materials. Around the same time, Proost and co-workers studied a series of Ni-Al compounds for efficient water oxidation. They synthesized a series of pure Ni, AlNi$_3$, AlNi, and a composite of AlNi$_3$ and Al$_2$Ni$_3$ as electrode material on a Si substrate and compared their activity towards OER. Out of all, AlNi$_3$ exhibited the highest activity with an $\eta_{10}$ of 300 mV operating stably at this current density under prolonged conditions for 4 h and a Tafel slope of 103 mV dec$^{-1}$. Additionally, annealing of as prepared Ni$_3$Al at different temperatures to study the coarsening effect on HER activated the electrocatalyst. The most stable and active species was received by annealing deposited Ni$_3$Al at 1160 °C. On the other hand, advanced stability, and high intrinsic activity were achieved by Yang’s group using a eutectic high entropy alloy (EHEA) as a template to synthesize the multicomponent porous structure (MCPS) FeCoNiCrNb$_6$ for OER catalysis. Their composite material showed excellent performance reaching 288 mV at $\eta_{10}$ with a Tafel slope of 27.67 mV dec$^{-1}$ and durability for 30 h.

Of late, earth-abundant and considerably acid-stable tantalum-based intermetallics have also been considered as promising candidates for OER in acidic media. Schaak and co-workers reported a bimetallic series of Ta with Co, Ni, and Fe were produced by arc melting from the elemental metal powders in the stoichiometric composition. The obtained intermetallic Ni$_2$Ta, Fe$_2$Ta, and Co$_2$Ta phases were pressed into pellets and used as electrodes for OER catalysis with Ni$_2$Ta showing the highest activity at an $\eta_{10}$ of 570 mV with the stability of over 60 h.

Transition metal silicides have gained much attention lately for their specific crystal and electronic structures with high conductivity and stability that are the decisive factors for catalytic water-splitting. For application in anion exchange membrane fuel cells, bifunctional cobalt and nickel silicide have been decorated on silicon-oxy-carbide (SiOC) and tested for OER and oxygen reduction reaction (ORR). The synthesized Co/SiOC contained Co$_2$Si and traces of pure Co while the Ni/SiOC consisted of a mixture of Ni$_2$Si and Ni$_5$Si. From both species, Ni/SiOC showed the highest performance in OER with $\eta_{10}$ of 390 mV and exhibited better OER kinetics during catalysis. Following this, Kumar et al. used the molten Li$_2$KI eutectic mixture to access NiSi and Ni$_5$Si via a reactive ionic Zintl solid (Na$_x$Si$_y$) and a Ni precursor. To reach a current density of 10 mA cm$^{-2}$ an $\eta$ of 570 mV was needed for NiSi and Ni$_5$Si, which was comparable to a commercial IrO$_2$ reference. In addition, porous Fe$_2$Si, Fe$_5$Si$_3$, and FeSi have been prepared as intermetallic anode for zinc electrowinning. While the Fe$_5$Si showed the best OER performance with $\eta_{10} = 955$ mV and a Tafel slope of 253 mV dec$^{-1}$, the Fe$_5$Si$_3$ ($\eta_{10} = 1072$ mV, Tafel slope = 327 mV dec$^{-1}$) exhibited better stability over 400 h.

FeNi$_3$ NPs on NC were synthesized by Mu and co-workers and the resulting FeNi$_3$@NC electrocatalyst displayed excellent OER activity of $\eta_{10} = 277$ mV (Tafel slope = 77 mV dec$^{-1}$) in alkaline conditions which they attribute to the unique structure with improved physical properties. To understand such systems more in-depth, Chen et al. studied a series of Fe-Ni intermetallics on NC gaining deeper insight into the role of iron in the structure. They synthesized various FeNi$_x$/NC, Fe–FeNi$_3$/NC, and Fe-enriched FeNi$_3$/NC electrocatalysts and found the Fe-enriched FeNi$_3$/NC was most active for OER with $\eta_{10}$ of 360 mV and a Tafel slope of 82 mV dec$^{-1}$ followed by Fe–FeNi$_5$/NC ($\eta_{10} = 390$ mV, Tafel slope = 86 mV dec$^{-1}$) and FeNi$_3$/NC ($\eta_{10} = 450$ mV, Tafel slope = 141 mV dec$^{-1}$) to reach the same current density concluding the higher Fe incorporation led to a positive effect giving more disorder to the structure. The influence of Fe in the Ni$_3$Al structure was also investigated by Bai et al., where the highest activity for OER was observed Ni$_{2/3}$Fe$_{1/3}$Al with an $\eta_{10}$ of 299 mV and a Tafel slope of 58.9 mV dec$^{-1}$, which was much lower compared to Ni$_{3/4}$Fe$_{2/4}$Al, Ni$_{5/6}$Fe$_{1/6}$Al, and other alloy materials.

Das and co-workers prepared ultra-small intermetallic NiZn phase using Ni NPs with organometallic Zn precursor using low-temperature solution chemistry. This non-precious electrocatalyst showed a substantial OER in alkaline solution with an estimated $\eta_{10}$ of 283 mV, a Tafel slope of 73 mV dec$^{-1}$, and stability of 16 h and surpassed the activity of pure Ni and Ni$_{5/6}$Zn$_{1/6}$ alloy. Similar observations were drawn also in a comparative study on NiSn coatings on electrodes used for alkaline OER. The electrodeposited NiSn films at different potentials showed higher intrinsic activity than bare Ni films.
and the most active intermetallic phase NiSn100 reached $\eta_{10}$ at 473 mV and a Tafel slope of 64 mV dec$^{-1}$.227

Antiperovskite materials are intermetallic compounds with a perovskite crystal structure but instead, the anion and cation positions are interchanged in the unit cell.228 Transition metals and nitrogen or carbon can form antiperovskite structures at suitable compositions (AXM$_3$: A = Cu, Al, Zn, etc.; X = N or C; M = Ni, Fe, Co, etc.), and can be well engineered to demonstrate versatile properties for OER catalysis.229 The p-Cu$_{1.5}$NNi$_{3.5}$/FeNiCu from Shao’s group exhibited superior OER activity ($\eta_{10}$ of 260 mV and Tafel slope of 52 mV dec$^{-1}$) in alkaline media with at long-term stability over 24 h exceeding the activity of IrO$_2$ (Fig. 6d–f).229 Inspired by this, Zou et al. developed a new intermetallic antiperovskite Co$_2$InCo$_{0.7}$Na$_{0.3}$ that showed impressive OER activity with $\eta_{10}$ of 260 mV, Tafel slope of 76.2 mV dec$^{-1}$ and moderate stability of 10 h, and presented valuable insights into the active structure of this unique material class.229

We investigated a manganese gallide (MnGa$_4$) and iron germanide (Fe$_x$Ge$_y$), both prepared by solid-state synthesis for potential application in alkaline OER. The MnGa$_4$ is a d-sp$^2$ bonded Hume–Rothery intermetallic compound with strong directional (covalent) bonds, metallic behavior, and antiferromagnetic ordering.78,219 The prepared MnGa$_4$ has a defective CsCl structure, where three-fourths of the Cs atoms are eliminated to form corner-linked cubes (MnGa$_{3.7}$) and proved to be a magnificent precatalyst for the electrochemical OER with a $\eta_{10}$ of 293 mV (Tafel slope = 98 mV dec$^{-1}$) for more than 24 h.219

Alternatively, Fe$_x$Ge$_y$ crystal structure consisted of a dense packaging of Fe and Ge atoms, built up by polyhedral of five different Fe atoms forming square pyramids, distorted octahedral and pentagonal prisms with the Ge atoms. In comparison to pure Fe-based materials, the Fe$_x$Ge$_y$ showed superior OER activity with $\eta_{10}$ of 221 mV and a Tafel slope of 32 mV dec$^{-1}$ and was constantly mediated the reaction under high potential for more than a day. In continuation, we further developed synthetic strategies to obtain NiGe$_{105}$ and FeAs$_{149}$ from molecular SSPs. The nanoparticle intermetallic phases proved to be very efficient during alkaline OER exhibiting low overpotentials and prolonged stability of three weeks.191 A chemical reduction approach was also utilized to derived phase pure FeSn$_2$ nanocrystals recorded one of the lowest $\eta_{10}$ of 197 mV (Tafel slope = 33 mV dec$^{-1}$) with the stability of over 60 h. This system was found to be even superior to analogous Fe-, FeNi as well as noble-metal-based materials.192 An up-to-date table on the activity parameters of intermetallics has been provided in Table 2.

The most active non-noble benchmarking systems for OER are based on metal oxides, (oxy)hydroxides, and chalcogenides. For instance, the $\eta_{10}$ of several present NiFe or NiCo (oxy) hydroxides and oxide systems range only between 180 to 240 mV with Tafel slopes of 28–35 mV dec$^{-1}$ that also remain stable for several hours even at higher current densities of 500 mA cm$^{-2}$ and 1000 mA cm$^{-2}$.21 Similarly, most of the other active materials such as perovskites, phosphides, alloys and carbides displaying overpotentials between 200–300 mV with Tafel slopes ranging from 40–80 mV dec$^{-1}$ to facilitate OER have lately been reported. Interestingly, the OER activities of these materials are matched closely with recently reported intermetallic phases and likely to improve significantly based on the structural tuning, composition, and electronic conductivity (see Section 8) making them an intriguing class of materials for further investigations.231

7. Intermetallics for overall water-splitting

Electrocatalytic overall water-splitting with intermetallic phases is a relatively nascent field with only a few reported examples.29,30 An early study on several alloys and intermetallics, e.g. Ti$_3$Ni, TiNi, TiCu, TiCo, and ZrNi from Miles revealed that proper combinations of elements can be favorably modified to facilitate HER and OER.

Although no further investigations on structure, morphology, or physical properties were carried out, a correlation on periodic trends of overpotential and atomic number with modern values for electronic work functions was derived for water splitting.29 To increase the roughness factor, Moller’s group fabricated AlNi electrodes for alkaline water electrolysis and highlighted the importance of large ESCA and low charge transfer resistance for efficient water splitting, although, the electrodes deteriorated dramatically during the harsh reaction conditions.229 Taking inspiration from the previous works, we carefully designed and developed a novel strategy for a controllable synthesis CoSn$_2$ nanocrystals that contained both active and conducting sites.231 As anticipated, CoSn$_2$ displayed an excellent $\eta_{10}$ of 230 mV for OER, $\eta_{10}$ of 103 mV, and necessitated a cell potential of 1.55 V to reach 10 mA cm$^{-2}$ in alkaline solution with almost 100% FE surpassing their Co and Sn counterparts.

In search of new overall water splitting electrocatalysts, Zhang and co-workers synthesized a noble metal-based intermetallic NiIrRuAl-1/3 with low content of Ir and Ru (34 at%) to form a hierarchically nanoporous nanowire structure with high ECSA that showed $\eta_{10}$ of 237 mV (Tafel slope = 50 mV dec$^{-1}$) for OER, while the NiIrRuAl-3/1 performed best in HER with $\eta_{10}$ of 14 mV (Tafel slope = 23 mV dec$^{-1}$) and a cell potential of 1.46 V was required for NiIrRuAl-1/3//NiIrRuAl-3/1 in an acidic media with extended stability of 35 h outperforming IrO$_2$ and Pt/C references.164 At the same time, a three-dimensional porous intermetallic MoNi$_4$ network was constructed by Jin et al. that demonstrated impressive performance as electrode material displaying $\eta_{10}$ of the only 28 mV (Tafel slope = 36 mV dec$^{-1}$) for HER, $\eta_{10}$ of 280 mV (Tafel slope = 79 mV dec$^{-1}$) for OER. When examined for bifunctional overall water splitting, a cell potential of 1.58 V at a current density of 10 mA cm$^{-2}$ was observed with over 24 h stability, which was significantly better than the reference Ni electrodes.123 Soon after, an Al$_3$(NiCoIrMo) intermetallic phase was accessed via solid-state route and partially dealloying to a reduced content of Ir in the structure (~20 at%). The quinary nanoporous np-AlNiCoIrMo HEA needed $\eta_{10}$ of 18.5 mV (Tafel slope = 33.2 mV dec$^{-1}$) for HER, $\eta_{10}$ of 233 mV (Tafel slope = 55.2 mV dec$^{-1}$) for OER, and overall cell voltage of 1.505 V with high durability of 48 h.169 Moreover,
Shi and co-workers developed self-supported monolithic nanoporous Co$_3$Mo/Cu electrodes from Cu$_{12-x}$Co$_x$Mo$_y$Al$_{88}$ precursors, which showed unexpected performance for HER with $\eta_{10}$ of merely 12 mV (Tafel slope = 40 mV dec$^{-1}$) while $\eta_{164}$ for OER was 350 mV (Tafel slope = 82 mV dec$^{-1}$) (Fig. 7). The catalyst was further tested for overall water-splitting.

### Table 2: Comparison of reported OER overpotentials of selected intermetallic catalysts at a given current densities in alkaline, acidic and neutral media$^a$

| Catalyst (substrate) | Electrolyte | Current density (mA cm$^{-2}$) | Overpotential (mV) | Tafel slope (mV dec$^{-1}$) | Reference |
|----------------------|-------------|-------------------------------|-------------------|----------------------------|-----------|
| FeAs (NF)            | 1 M KOH     | 10                            | 252               | 32                         | 149       |
| FeSn$_2$ (NF)        | 1 M KOH     | 10                            | 197               | 33                         | 102       |
| Co$_3$Mo (direct)    | 1 M KOH     | 10                            | 230               | —                          | 221       |
| Fe$_6$Ge$_5$ (FTO)   | 1 M KOH     | 10                            | 299               | 89                         | 221       |
| Fe$_3$Ge$_5$ (NF)    | 1 M KOH     | 10                            | 228               | 56                         | 98        |
| NiGe (FTO)           | 1 M KOH     | 10                            | 322               | —                          | 105       |
| NiGe (NF)            | 1 M KOH     | 10                            | 228               | 56                         | 105       |
| Porous MoNi$_4$ networks (Ni)| 1 M KOH | 10                        | 280               | 79                         | 131       |
| Co$_3$Mo (direct)    | 1 M KOH     | 164                           | 350               | 82                         | 113       |
| FeNi$_{0.3}$@NC (GC) | 1 M KOH     | 10                            | 277               | 77                         | 225       |
| FeNi$_{0.3}$@NGC (GC) | 0.1 M KOH     | 10                           | 450               | 141                        | 103       |
| Fe-enriched FeNi$_{0.3}$@NGC (GC) | 0.1 M KOH | 10                        | 360               | 82                         | 103       |
| MnGa$_3$ (NF)        | 1 M KOH     | 10                            | 293               | 98                         | 97        |
| Co/SiO$_2$@GC (GC)   | 0.1 M KOH   | 10                            | 440               | 118                        | 223       |
| Ni/SiO$_2$@GC (GC)   | 0.1 M KOH   | 10                            | 390               | 90                         | 223       |
| Ni$_3$Si (GC)        | 0.1 M KOH   | 10                            | 570               | 119                        | 224       |
| Ni$_3$Si (GC)        | 0.1 M KOH   | 10                            | 570               | 94                         | 224       |
| Ni$_3$Ta (direct)    | 0.5 M H$_2$SO$_4$ | 10               | 770               | —                           | 110       |
| Co$_3$Ta (direct)    | 0.5 M H$_2$SO$_4$ | 10               | 600               | —                           | 110       |
| Fe$_3$Ta (direct)    | 0.5 M H$_2$SO$_4$ | 10               | 770               | —                           | 110       |
| IrNi-RF (GC)         | 0.1 M HClO$_4$ | 10               | 313.6             | 48.6                        | 120       |
| FeSi (direct)        | 1.6 M H$_2$SO$_4$ | 10              | 1107              | 634                         | 90        |
| Fe$_3$Si (direct)    | 1.6 M H$_2$SO$_4$ | 10              | 1072              | 327                         | 90        |
| Fe$_3$Si (direct)    | 1.6 M H$_2$SO$_4$ | 10              | 953               | 253                         | 90        |
| PtCo$_{1.2}$@GC (GC) | 1 M KOH     | 10                            | 380               | 71.2                        | 167       |
| PtCo$_{1.2}$@GC (GC) | 1 M KOH     | 10                            | 386               | 65.9                        | 167       |
| IrRu@Te (GC)         | 0.5 M H$_2$SO$_4$ | 10               | 220               | 35                          | 132       |
| IrRu@Te (GC)         | 0.5 M H$_2$SO$_4$ | 10               | 245               | —                           | 132       |
| IrRu@Te (GC)         | PBS         | 10                            | 309               | —                           | 132       |
| Al$_2$Pt (direct)    | 0.1 M HClO$_4$ | 10              | 450               | —                           | 108       |
| AlNi$_3$ (Si)        | 1 M KOH     | 10                            | 300               | 46                          | 219       |
| Al$_{2}$Ni$_{3}$/Al$_{12}$Ni$_{3}$ (Si) | 1 M KOH | 10                        | 390               | 52                          | 219       |
| AlNi$_3$ (Si)        | 1 M KOH     | 10                            | 310               | 47                          | 219       |
| Ni$_3$Al N-0 (direct) | 1 M KOH | 10                      | 320               | 112                         | 95        |
| Ni$_3$Al N-1160 (direct) | 1 M KOH | 10                      | 280               | 73                          | 95        |
| Ni$_3$Al N-1200 (direct) | 1 M KOH | 10                      | 293               | 85                          | 95        |
| Ni$_3$Al N-1240 (direct) | 1 M KOH | 10                      | 300               | 103                         | 95        |
| Ni$_3$Al N-1280 (direct) | 1 M KOH | 10                      | 305               | 108                         | 95        |
| Ni$_3$Fe$_{1.2}$ (direct) | 1 M KOH | 10                      | 299               | 58.9                        | 169       |
| Co$_3$InCo$_{0.7}$N$_{0.3}$ (GC) | 1 M KOH | 10                      | 350               | 76                          | 229       |
| CuNNi$_{1.4}$+Cu (GC) | 1 M KOH     | 10                            | 510               | 163                         | 99        |
| CuNNi$_{1.2}$ (GC)   | 1 M KOH     | 10                            | 480               | 136                         | 99        |
| Cu$_{1.2}$NNi$_{1.8}$/FeNiCu (GC) | 1 M KOH | 10                      | 300               | 76                          | 99        |
| p-Cu$_{1.8}$NNi$_{1.2}$/FeNiCu (GC) | 1 M KOH | 10                      | 280               | 52                          | 99        |
| NiZn (GC)            | 1 M KOH     | 10                            | 283               | 73                          | 226       |
| Cu$_3$Sn@Cu (Cu)     | 1 M KOH     | 50                            | 384               | 177                         | 232       |
| NiSn$_{10}$ (Ni)     | 1 M NaOH    | 40                            | 541               | 60                          | 227       |
| NiSn$_{60}$ (Ni)     | 1 M NaOH    | 40                            | 516               | 63                          | 227       |
| NiSn$_{100}$ (Ni)    | 1 M NaOH    | 40                            | 473               | 57                          | 227       |
| FeCoNiCrNb$_{0.5}$ (direct) | 0.1 M KOH | 10                      | 288               | 27.67                        | 101       |
| AlNiHT10 + L (Ni)    | 1 M KOH     | 10                            | 338               | —                           | 160       |

$^a$ FTO = fluorine doped tin oxide, GC = glassy carbon, NF = nickel foam.
resulting in a cell potential of 1.65 V to produce a current density of 100 mA cm$^{-2}$.

It should be noted here that the activity presented in Tables 1 and 2 are only based on the overpotential, Tafel slope, electrolyte, and substrate effects. As the experimental conditions (pH, temperature, loading amount, film thickness, $iR$ compensation, etc.) can differ in each case, the reader is referred to the respective references.

8. Insights into the active structures

Intermetallics are considered as a highly stable compound with higher resistance against chemical oxidation than single metal compounds due to their ordered structure with d-orbital interaction and covalent bonding. Currently, the high activity of intermetallic phases in the two different half-reactions of water splitting is often ascribed to the pronounced synergism of the different metallic species (active or electrically conductive), when no change in the original or pristine state of the structure was observed under catalytic conditions. If the catalyst is not stable during OER or HER catalysis, then it will be in its precatalytic state and the so-called pre-catalyst can transform in three possible ways: (i) near-surface, (ii) partial (core–shell), and (iii) complete transformation resulting into crystalline or amorphous structures (Fig. 8). In the case of near-surface, only the outermost layer of the catalysts are transformed typically by leaching or oxidation so that the reactants can participate in the reaction and contribute towards the net catalytic activity. The extent of transformation mainly depends upon the porosity or voids created by leaching (non)metals. In many cases, the structural transformation of the precatalyst is not continuous and ceased after a certain time interval when electrolyte can no longer penetrate through the precatalyst’s core forming a stable core–shell phase. Under prolonged electrolysis, this process encompasses deep inside the core of the particles finally forming a stable active structure. The degree of conversion is not only limited to the leaching phenomenon but also depends on the size, morphology, composition of particles as well as the complex bonding situations of intermetallics. The transformation primarily results in increased ECSA, enhanced defect/disordered sites, higher electrical conductivity, surface or bulk activity, and structural porosity. In the following, we describe the type of transformation attained for intermetallic phases under electrocatalytic HER and OER conditions.

8.1 Active sites in HER

Most of the reported intermetallic compounds have shown no indication of transformation acting as chemically stable and highly resistant in a harsh HER environment. The high activity is explained by the specific surface composition of the...
formed materials.\textsuperscript{48} For instance, the remarkable HER performance of Ni and rare-earth-based intermetallics (Ni–LaNi\textsubscript{5} and Ni–MnNi\textsubscript{5–x}Co\textsubscript{x}Al\textsubscript{2}Al\textsubscript{2}) was discussed in light of Brewer–Engel valence bond theory implying a synergistic effect of the hyper-electronic character of the Ni and the hypo-electronic character of the rare-earth element on the electrode surface to facilitate H\textsuperscript{+} adsorption (Heyrovsky step).\textsuperscript{222} The same synergistic effect of intermetallics was also revealed for Ni\textsubscript{1.5}Mo and Ni\textsubscript{2}Mo where d-orbitals of nickel (group VII) are more filled than those of molybdenum (group VI). Upon alloying, the d-orbitals interact and the catalyst becomes more active for HER when the d-band vacancies are introduced and becomes less active as d-band vacancies are filled.\textsuperscript{49} Similar observations have also been made for PtNi, LaNi\textsubscript{5}, TiNi\textsubscript{3}, ZrNi\textsubscript{3}, HfPd\textsubscript{3}, MoPt\textsubscript{3}, FeR (R = rare earth metals), and Ni–Mo–NGTs.\textsuperscript{99,104,118,176,222} In the case of NiIrRuAl, the charge transfer induced Ni gave rise to modified surface valence states that optimized the H\textsuperscript{+} adsorption and improved the catalytic HER activity.\textsuperscript{24} Pt\textsubscript{5}Si showed higher catalytic HER activity than Pt, by facilitating HO–H bond breaking and H\textsuperscript{+} adsorption through the incorporated Si and remained unchanged in the catalytic process.\textsuperscript{226} In addition, Pt\textsubscript{5}Ti\textsuperscript{139} and TiCO\textsubscript{2} (ref. 109) also showed ideal conditions to adsorb H\textsuperscript{+} on the surface. Besides the optimal adsorption of H\textsuperscript{+} (Fig. 9c), the Pt\textsubscript{5}Ti proved to be very stable under electrocatalytic conditions in acidic HER. Only a minor change in activity was measured after 2000 CV cycles and the TEM showed that Pt\textsubscript{5}Ti phase was still intact (Fig. 9a and b).\textsuperscript{199}

Along with the synergistic effect to enhance structural and electronic properties, and enhancement of the mass-specific surface area in the intermetallic phases has been observed, which significantly contributes to the HER activity.\textsuperscript{224,157,143} NiMo exhibited a much higher surface area and a larger turnover frequency per-surface atom resulting in a higher mass activity for HER. The tested intermetallic phase was stable in alkaline media and only degraded in acidic media. By the calculations of DFT and Bader charge population, it was demonstrated that NiMo had its optimal adsorption conditions and was stable when NiMo was covered with holey graphene and therefore possessed more active sites (Fig. 9d and e).\textsuperscript{206} By additionally covering the surface of NiMo with graphene, the dissolution of the intermetallic phase in acidic media was drastically reduced and enhanced the catalytic activity of the material.\textsuperscript{130} Ni\textsubscript{20}Mo\textsubscript{40}Ti\textsubscript{10} and Ni\textsubscript{30}Mo\textsubscript{50} were also found to be stable and highly active due to their enlarged surface area. Additionally, it was found that the presence of Ti in NiMo further increased the intrinsic activity for HER.\textsuperscript{168} The comparably high surface area that maximizes the exposure of active sites was also observed in Ti–Ni,\textsuperscript{206} Fe–Zn\textsuperscript{155} Pd\textsubscript{3}Si,\textsuperscript{225} Ni\textsubscript{2}Si,\textsuperscript{155} GaPt\textsubscript{5},\textsuperscript{122} GaPd\textsubscript{3},\textsuperscript{123} and Pt–Dy\textsuperscript{67} resulting in high HER performance. Jin et al. ascribed their efficiency and durability for their np-AlNiCoIrMo HEA to the high-entropy stabilizing effect of the quinary compound. The five different uniformly distributed elements with unique physiochemical properties attributed synergistically to the formation of the porous structure that was retained during HER.\textsuperscript{108} Similar conclusions were drawn for the FeCoNiAlTi HEI.\textsuperscript{162} Intriguingly, the bifunctional Co\textsubscript{3}Mo/Cu exhibited efficient HER and remained unchanged. The high activity of Co\textsubscript{3}Mo was attributed to the surfaces precisely composed of Co and Mo atoms, wherein their dissimilarity results in both ligand effect and strain effect to regulate the adsorption energies of the H* intermediate, leading to a high performance HER electrode.\textsuperscript{113}

Apart from electrochemically stable electrodes, many reports show surface passivation and even transformation of the precatalyst during HER. In such cases, the transformation of the surface of the film or particles leads to a core–shell structure and the active materials typically exhibit an increased surface area.\textsuperscript{234} Despite reports on electrochemically stable Ni\textsubscript{3}Al electrodes for HER, Al-based electrodes tend to transform during catalysis due to Al leaching.\textsuperscript{93,94,235} Highly porous Ni\textsubscript{Al} from Liu’s group showed slight surface passivation after strongly alkaline (6 M KOH) HER catalysis. It was expected to form NiO and NiOOH/OOH on the surface, but it is unclear whether the formed species influenced the HER activity.\textsuperscript{231} In other reports, the Al leaching in alkaline solution is intended to increase the electrode activity by increasing the surface (porosity effect) through the created voids, and some phases were not stable and disintegrate during leaching.\textsuperscript{226} However, the intensity of the leaching process was dependent strongly on the initial stoichiometric and elemental composition, the structure, and the morphology of the starting material as well as the electrolyte.\textsuperscript{234} A loss of Al was also confirmed for Al–Cu\textsubscript{4}Ni@Cu\textsubscript{4}Ni in alkaline HER forming a core–shell structure. The stability of the material was ascribed to the stable Cu\textsubscript{4}Ni shell, in which the Ni atoms anchored with subsurface Al atoms forming strong Ni–Al bonds and suppress diffusion of Cu atoms.\textsuperscript{464} The same results were obtained for PdCu\textsubscript{3} and PdBi\textsubscript{2} where a loss of Cu and Bi resulted under HER forming in situ a Pd rich catalyst. Long-term HER experiments confirmed that the shell of the catalyst inhibited further leaching and remained stable once formed that indeed increased the H\textsuperscript{+} adsorption as well as the charge transfer through the catalyst.\textsuperscript{119,129} Analogous HER reaction pathways
were observed when we investigated CoSn$_2$ as a catalyst for water splitting. The surface leaching of Sn from the outermost layer of CoSn$_2$ exposed abundant Co$^0$ sites for HER for facile H$^+$ adsorption and reduction to form H$_2$ while unchanged bulk phase contributed in terms of electronic conductivity.

A more drastic transformation of an intermetallic phase into an oxide phase during HER was observed in Ni–Co–Al on a Ni-foil by Zhou et al. X-ray photoelectron spectroscopy (XPS) data clearly revealed, that after HER, Al was oxidized into Al$_2$O$_3$; Ni into NiO and Al$_2$NiO$_4$ and Co into Co$_3$O$_4$ and Al$_2$CoO$_4$. Since no oxide species was present on the precatalyst material, the Al$_2$NiO$_4$/Al$_2$CoO$_4$ was considered as an active species contributing to the electrocatalytic performance towards HER in alkaline solution.

8.2 Active sites in OER

In stark contrast to HER catalysts, the strongly oxidizing conditions of the OER force most of the non-oxides inexorably to form oxides and (oxy)hydroxides (at least at the surface) due to their thermodynamically more favorable stability. However, some of the reported intermetallic compounds show almost no or limited change in their crystal structure and morphology under OER conditions. For example, Mukherjee and co-workers reported intermetallic PtCo with CoO$_x$ composite for efficient OER and ORR. The sponge-shaped CoO$_x$ matrix surrounding PtCo NPs enforced synergic “spillover” effects and helped to preserve the structural and morphological integrity of PtCo NPs preventing their aggregation and dissolution in alkaline electrolytes. Likewise, the microstructure and element distribution of the np-AlNiCoIrMo HEA revealed no obvious microstructure coarsening and uniformly element distribution without aggregation after the OER durability test. It was stated that the bicontinuous nanoscale ligament-pore structure facilitates mass transport (both electrolyte and gas) and provided a conductive network for fast electron transfer and efficient electrochemical reactions. No visible transformation was reported for the intermetallic Ni/SiOC phase even after 10 000 CV cycles of OER which was attributed to the presence of Ni$^{III}$, giving increased conductivity to the material and providing higher affinity towards adsorption of OH$^-$ anions enhancing the OER. Retention of structural stability after OER was also reported for intermetallic AlNi$_3$, AlNi$_3$/Al$_2$Ni$_5$, and AlNi phases, however, a minor amount of Al leaching was unavoidable. As discussed earlier, in most cases, the transformation of the (pre)catalyst into the active species during OER catalysis is imminent. The nature of the formed catalyst is mainly defined by the structure of the precatalyst, electronic properties, the leaching ability and size of the leaching ion, and the
transformation conditions strongly influenced by the pH of the electrolyte as well as the applied potential. The intermetallic NiSn synthesized by Jović et al. slowly oxidized on the surface into a β-NiOOH in alkaline OER that γ-NiOOH activated from bare Ni. Using Raman spectroscopy, Shen’s group was also able to uncover in situ NiOOH active layer by etching of MoNi₅ in alkaline OER conditions. The core–shell Cu₁₋₂NNi₃₋₃FeNiCu, reported by Shao’s group described the FeNiCu-(oxy) hydroxide as the active species in the HER process, that retained structural integrity post 24 h of OER. The group of Grin observed that the bulk Hf₂B₂Ir₅ and major volume close to the surface did not change under the OER conditions. The formation of monoclinic Hfo₂ and IrO₂(OH)₆(SO₄)₂ particles upon anodic treatment of the Hf₂B₂Ir₅ was mainly related to the oxidation of the secondary phase HfB₂Ir₃ that was also present in minor amounts in addition to the self-controlled near-surface oxidation of Hf₂B₂Ir₅. They concluded that the chemical bonding features of the Hf₂B₂Ir₅ compound with a cage-like Ir–B framework, hosting Ir atoms, only allow near-surface oxidation and inhibit deep Ir leaching. Zhang and co-workers showed that the surface oxidation of NiRulAl led to an improvement of catalytic activity. By tuning the atomic ratio of Ir/Ru, it was demonstrated that a higher Ru content improves the intrinsic catalytic activity while more Ir effectively generates Ir oxides on the surface to retain the catalytic stability. More Ni content weakened the adsorption of oxygen-based intermediates by shifting down the Ir d-band center signifying its role in OER activity and it also leached from the surface structure leading to an increased concentration of oxygen species, drastically enhancing the OER activity. The persistent surface oxidation of intermetallics under OER often leads to a core–shell structure that usually consists of a crystalline or amorphously active shell and a conductive core that supports cooperative interactions of the elements. While thickness and properties of the surrounding layer vary depending on the starting material and transformation conditions, it is often observed that the oxide coating actively inhibits further oxidation of the material underneath. As an example, Ni₃Ta, Co₃Ta, and Fe₃Ta anodes slowly corroded under acidic OER conditions and formed a protective oxide layer that indeed acted as a passivating oxide coating, allowing subsurface atoms to continue to participate in OER while limiting their dissolution. In another example, acid etching of IrNi to obtain nanocages induced an IrO₃ layer on the surface. This layer increased under continuous OER operation resulting in core–shell IrNi@IrO₃. The unique feature of the inner lying IrNi and outer IrO₃ phase was concluded to be responsible for the high OER activity. In a recent approach, we observed similar transformations in FeSn₂. After alkaline OER, the FeSn₂ generated a goethite α-FeOOH shell with FeSn₂ core. The α-FeOOH shell facilitated the OER while the intact intermetallic core enhanced electrical conductivity. Besides, the Ni₂Si and NiSi prepared by Kumar et al. suffered substantial reorganization of the structure for alkaline OER. The SiO₂ species leached from the surface continuously to form an active NiOOH shell on both NiSi or Ni₂Si. In many cases, the complete degradation of the precatalyst into a more crystalline or amorphous (oxy)hydroxide under OER has been reported. As the material is oxidized, it undergoes significant leaching of the nonmetallic component and facilitates major structural reorganization with substantially enhanced ECSA from the newly porous bulk-active structures that in turn increases the number of active sites for the OER. Different observations were made by Møller’s group for their Al–Ni phase where they postulated that in alkaline conditions the material is thermodynamically not stable at zero potential and undergoes corrosion. Another possibility for dissolution was localized acid formation inside the porous structure during OER resulting from disturbances of oxygen bubbles on OH⁻ migration into the inner active sites of the catalyst. As a consequence the pH is locally decreased, hence the structure is further weakened and corrosion will ultimately severely damage the structure. Similar observations were also made for intermetallic MnGa₄ where three distinct crystalline MnO₇ phases: birnessite δ-MnO₂, leiptknechtite β-MnOOH, and hausmannite α-Mn₁₃O₄ were confirmed after the alkaline OER that was accompanied by an almost complete Ga loss from the structure (Fig. 10). The transformation trend was also extended for SSP-derived FeAs and NiGe precatalysts. In the case of FeAs, an active 2-line ferrihydrite phase was formed after alkaline OER with the severe dissolution of As in the structure. The Raman and in situ XAS described the reaction pathway and the defective edges/sites, as well as the presence of surface tetrahedral coordinated FeIII atoms, were ought to be the active centers for OER catalysis. On the other hand, NiGe produced γ-NiOOH. The activity was attributed to the structural flexibility of Ni sites triggered by the defective structure with ionic intercalation of OH⁻/CO₃²⁻ between the large interplanar spacing of γ-NiOOH. In both cases, the transformation resulted in higher ECSA values, and better electronic conductivities to promote favorable OER kinetics with improved charge transfer properties to facilitate the OER. Of late, we reported an intermetallic Fe₉Ge₃, as a novel alkaline OER precatalyst forming first an in situ metastable core–shell structure that slowly collapses in prolonged OER conditions finally to form a FeIII(OH)₅H₂O. We recently investigated CoSn₂ for overall water splitting where after alkaline OER, CoSn₂ underwent a slow but rather complete structural transformation (Fig. 11). The loss of Sn from the structure of CoSn₂ led to a disordered, defected, and vacant amorphous active material with a CoO₉/CoOOH phase (with CoIII centered active species) that enables optimal adsorption of the oxygen species (Oₐd), thus facilitating the formation of adsorbed -OOH species (OOHₐd) by a nucleophilic attack, thereby promoting the deprotonation of OOHₐd to produce O₂. However, in the case of HER, only surface Sn loss was observed exposing the Co⁹⁺ site for H⁺ adsorption (Fig. 11). In all cases, the stability of the catalysts was also linked to the extent of structural transformations. If the transformation is very rapid and complete, no change in the stability curve is observed as the formed species are already sufficient to withhold the stable current under longer run. However, if the transformation is slower, an initial decrease in current was observed until a formation of core–shell is attained (not allowing further electrolyte permeability) and then a constant current at applied potentials is achieved. In both cases, the extent of
transformation was found to be important to produce an enormous number of active sites, increased surface areas, or even electronic conductivity.

9. Conclusion and challenges

Intermetallic compounds have emerged as compelling advanced energy materials for HER, OER, and overall water splitting. In various examples, intermetallic catalysts are displayed to promote the challenging reactions of water splitting with enhanced activity, improved reaction kinetics, and long-lasting durability. In this perspective, we have highlighted the suitability of well-defined intermetallic (pre)catalysts for the design of high-performance electrocatalysts with complex interface structure, bonding characteristics, and electronic properties, which is essential to increase overall energetic efficiencies of the water-splitting reaction and stabilization of their active structures under operating conditions. Although

Fig. 10 Proposed mechanism for the electroconversion of MnGa₄ to δ-MnO₂ with its intermediates (top). TEM images of MnGa₄ and the transformed oxide species after OER in alkaline media (bottom left). EDX mapping of Mn and Ga in MnGa₄ before alkaline OER and Mn and O mapping after. Adapted with permission from ref. 97, copyright 2019, Wiley-VCH.

Fig. 11 A synthetic approach to nanostructured CoSn₂, with structural modification during alkaline HER and OER (left). TEM images of showing amorphous Co₃O₄/CoSn₂ after OER (top right) and crystalline CoSn₂ after HER (bottom right) with SAED inset, scale bar = 10 nm. Adapted with permission from ref. 221 copyright 2018, Wiley-VCH.
substantial efforts have been devoted to exploring intermetallic compounds as novel catalysts for water splitting, they are relatively underdeveloped. Therefore, several critical aspects need to be taken into account for the further development of this field and are discussed below.

One of the essential factors for designing an efficient electrocatalyst is by synthesizing the materials in nanoscale, increasing the number of active sites, active surface areas, and electronic conductivities. The properties of intermetallic compounds are closely associated with their size, shape, structure, composition, and crystal phase and hence, they must be tuned and optimized according to the need of electrocatalysis. In order to form an intermetallic compound, a large amount of energy is required, which is usually achieved with high-temperature annealing methods. Such solid-state techniques often predictably give rise to large agglomerated particles or even by-products and in most cases, with significantly decreased catalytic active surface areas. Thus, it is extremely interesting to design low-temperature novel synthetic strategies that can allow better atomic-level control over the stoichiometry, high dispersity of nanostructures with access to unique electronic and surface structures. In this regard, low-cost techniques such as wet impregnation, SSP, chemical reduction, and electrochemical approaches entail more attention. Similarly, looking at the various known classes of intermetallic compounds with different element combinations, a knowledge-guided simulation could also be highly important for their rational synthesis with desired HER and OER properties.

Most importantly, it is now well-known that most of the intermetallic catalysts transform (either step-wise or completely) from their pristine state to active state under catalytic operating HER and OER conditions. Therefore, understanding the dynamic behavior of the catalyst through advanced ex situ and operando techniques to reveal the real active species, morphological and electronic changes, surface/bulk structure, and structure–activity relation during catalysis is of utmost significance. Indeed, efforts should also be devoted to testing the active intermetallic catalysts for OER and HER for a longer period at higher currents to observe their chemical stability. This will not only will help us to understand the reaction mechanism but also to further optimize the catalyst design with desired elements.

As the current studies have demonstrated the applicability of intermetallic compounds for electrocatalytic water splitting, it is now essential to examine them (either unfunctional or bifunctional) in industrially relevant conditions at elevated temperatures as well as at higher current densities in harsh alkaline or acidic environments. This step is pivotal, as the reaction conditions set for the lab-scale are entirely different when compared to the actual working electrolyzers. In addition to this, emphasis should also be given to the stability and the degradation of catalysts under operating conditions. Thus, future studies should be focussed on designing nanostructured intermetallic electrocatalysts with specific structure-types with porosity, precise control in their particle size and morphology with higher surface areas, enhanced electronic conductivity, structures with electrolyte permeability and bulk-activity, fast-redox switching sites, the maximum amount of edge sites, which have already been proven beneficial for non-oxidic materials. Like-wise, such designed intermetallics are expected to work under high current densities, elevated temperature, all-pH electrolytes, and for seawater splitting. In the near future, the research should also be concentrated on producing large electrodes with excellent mechanical and (electro)chemical stability, resistance to acidic/alkaline media, and bifunctionality.

The tunability of intermetallic materials by varying different metals could make this class of materials also interesting for electrocatalytic applications beyond water-splitting. In this context, we recently combined the electrochemical water-splitting and selective (almost 100%) oxygenation of organic substrates through in situ surface modification of intermetallic FeSn₂ precatalyst. Selective oxygenation is a demanding approach and bears higher economic value than oxygen produced by water splitting. Therefore, intermetallic catalysts could be used to further explore this nascent field where H₂ and oxygenated products can be liberated simultaneously, making it a lucrative technology. Alternatively, the same electrochemical technology with intermetallics can be applied in non-aqueous solvents to drive the other essential transformations such as regeneration of the triphenylphosphine from triphenylphosphine oxide as well as a one-pot Wittig olefination reaction that presently best driven by noble-metal catalysts.

Furthermore, the scope of the intermetallics can easily be extended electrochemical carbon dioxide (CO₂) reduction, a promising reaction to mitigate CO₂ emissions, where presently transition metals (in particularly Cu) have been used predominantly to yield high-end multicarbon products. Notably, the intermetallics can also be of special interest for the kinetically sluggish oxygen reduction reaction (ORR), an important cathodic reaction of fuel cell which is presently mediated by the metals, alloys, and carbons, etc. Another alternative strategy would be to utilize intermetallics to achieve a systematic electrochemical reduction of heavy non-metal oxides with very strong E–O Bonds (E = Si, P, S). Besides, intermetallics have also shown promising electrocatalytic behavior methanol oxidation reaction, surface coatings, and supercapacitors. Although intermetallics is relatively an unexplored field for electrocatalysis, more fundamental and applied research pursued simultaneously to uncover novel classes of materials and study their intriguing unusual properties toward energy applications.

Author contributions

C.W. conducted the literature research and wrote the first draft of the manuscript. P.W. M. initiated the overall idea and contributed to the planning and writing of the manuscript. M. D. and P.W. M. discussed, supervised, and edited the final version of the perspective.

Conflicts of interest

There are no conflicts to declare.
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