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Perovskite and layered perovskite oxynitrides for efficient sunlight-driven artificial synthesis

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Perovskite and layered perovskite oxynitrides are regarded as promising visible-light-responsive semiconductors for efficient artificial photosynthesis to produce renewable value-added energy resources, including H₂, formic acid (HCOOH), and ammonia (NH₃). This is because of their chemical stability in aqueous electrolytes at various pH values, along with a narrow bandgap and a suitable band position bestowing ideal optical properties. In this review, we discuss recent advances in photocatalysis using perovskite and layered perovskite oxynitrides responsive to a wide wavelength range of the visible-light spectrum. Next, we address in detail how the photoactivity of oxynitrides can be enhanced with respect to their synthesis, including bulk and surface engineering such as doping (or substitution), controlling their morphology, and crystal facet engineering. Finally, we discuss the existing challenges to realizing efficient artificial photosynthesis using these materials.

1. Perovskite and layered perovskite oxynitrides

Perovskite-type oxynitrides are mixed metallic compounds with the general formula AB(O,N)₃, where A and B represent a large alkali-earth metal or La (*i.e.*, A = Ca, Sr, Ba, or La) and a relatively small transition metal (*i.e.*, B = Ti, Ta, or Nb), respectively.^{1–3} The stoichiometry of anions O and N in the perovskite is determined by the combination of cations at the A and B sites. Fig. 1 displays the crystal structure of an ideal cubic

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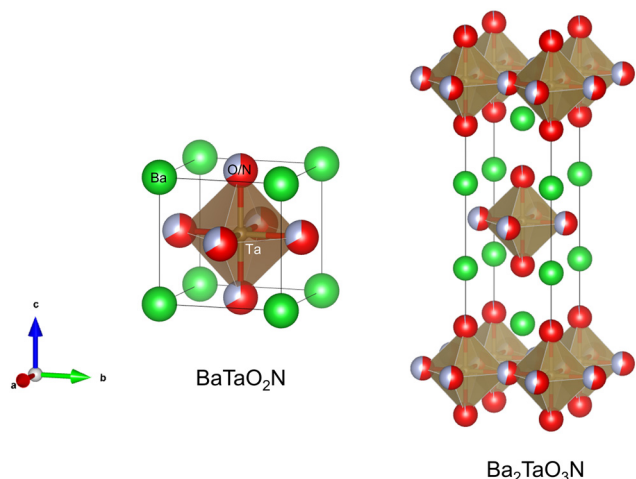


Fig. 1 Crystal structures of perovskite oxynitride BaTaO_2N and Ruddlesden-Popper (RP) phase layered perovskite oxynitride $\text{Ba}_2\text{TaO}_3\text{N}$.

perovskite $\text{AB}(\text{O},\text{N})_3$ whereby the $\text{B}(\text{O},\text{N})_6$ octahedron is at the center of the cube while the large A atoms occupy the corners. The relationship between the ionic radii (r) expressed as $r_{\text{A}} + r_{(\text{O},\text{N})} = \sqrt{2}(r_{\text{B}} + r_{(\text{O},\text{N})})$ is valid for the ideal cubic perovskite structure. The ratio of left- to right-hand sides of the expression is called the tolerance factor t , and the ideal cubic perovskites have a t value of unity. Tilting of the octahedron in the cubic structure originates from the subtle differences between the ionic radii of the cations leading to a t value less than unity. However, structural distortion producing orthorhombic or tetragonal shapes is commonly found, except for very few combinations including BaBO_2N ($\text{B} = \text{Ta}$ or Nb). In addition, the perovskite oxynitrides can adopt the $\text{A}_{1-x-1}\text{A}_{2-x}$ and/or $\text{B}_{1-y-1}\text{B}_{2-y}$ configurations at the A and/or B sites. In these cases, the stoichiometry of the anions deviates from the ideal coordination configuration if the cations at the A and B sites undergo a change in their oxidation state.

Besides the ideal cubic structure, perovskites can also adopt layered structures in which $\text{AB}(\text{O},\text{N})_3$ layers are separated by thin layers of intrusive materials (commonly metal oxides), as shown in Fig. 1. Depending on the intruding layers, the perovskite derivatives designated layered perovskites are generally classified as (i) Aurivillius ($(\text{Bi}_2\text{O}_2)(\text{A}_{n-1}\text{B}_n\text{O}_{3n+1})$), (ii) Ruddlesden-Popper (RP) ($\text{A}_{n+1}\text{B}_n\text{O}_{3n+1}$ or $\text{A}'\text{A}_{n-1}\text{B}_n\text{O}_{3n+1}$), or (iii) Dion-Jacobson ($\text{A}'\text{A}_{n-1}\text{B}_n\text{O}_{3n+1}$) phases ($n = 1, 2$, or 3 represents the number of stacked octahedra separated by a rock salt AO layer). The oxygen in the layered perovskites can be partially exchanged with another anion such as halogen and nitrogen. Sillen-Aurivillius layered perovskite oxyhalides $\text{A}_4\text{A}'_{n-1}\text{B}_n\text{O}_{3n+5}\text{X}$ (A and $\text{A}' = \text{Sr}, \text{Ba}, \text{Pb}$, or Bi ; $\text{B} = \text{Ti}, \text{Nb}$, or Ta ; $\text{X} = \text{Cl}$ or Br) were prepared by the substitution of oxygen by halogen.^{4,5} Moreover, RP phase oxynitrides $\text{A}_{n+1}\text{B}_n(\text{O},\text{N})_{3n+1}$ and Dion-Jacobson phase oxynitrides $\text{A}'\text{A}_{n-1}\text{B}_n(\text{O},\text{N})_{3n+1}$ ($\text{A} = \text{Ca}, \text{Sr}, \text{Ba}$, or La ; $\text{A}' = \text{Li}, \text{Na}, \text{K}$, or Cs ; $\text{B} = \text{Ti}, \text{Nb}$, or Ta), composed of mixed anions of O^{2-} and N^{3-} , have been studied as a novel layered perovskite oxynitride family active for various artificial photosynthesis approaches.^{6–10}

Perovskite oxides (e.g., SrTiO_3) and layered perovskite oxides (e.g., $\text{Ba}_5\text{Nb}_4\text{O}_{15}$) (well-known active catalysts for water splitting to produce H_2 gas) possess large bandgap energy values (E_{g}) of up to or even above 3.2 eV ($\lambda < 380$ nm), which are too high to harvest a wide range of the solar spectrum, including visible light.^{11–13} Introducing N into the oxide (i.e., nitridation) shifts the valence band maximum (VBM) to a more negative potential because the potential of the N 2p atomic orbitals is lower than that of the O 2p orbitals.¹⁴ Although nitridation leaves the conduction band (CB) potential of the oxide almost unchanged, hybridization of the N 2p and O 2p orbitals during nitridation narrows the E_{g} value of the oxide, resulting in a visible-light-responsive perovskite or layered perovskite oxynitride.

The wavelength edges of the visible-light absorption by perovskite oxynitrides vary depending on the combination of cations at the A and B sites, as presented in Fig. 2. For instance, SrNbO_2N can be prepared by the partial exchange of 3O^{2-} for 2N^{3-} during the nitridation of layered perovskite $\text{Sr}_5\text{Nb}_4\text{O}_{15}$, with the resulting reddish-brown oxynitride SrNbO_2N able to absorb visible light up to a wavelength of approximately 680 nm.¹⁵ In the same manner, in the syntheses of SrTaO_2N , LaTiO_2N , BaTaO_2N , and BaNbO_2N , complete nitridation of the corresponding starting oxide resulted in oxynitrides capable of absorbing photons up to ca. 560, 620, 660, and 740 nm, respectively.^{16–19} The apparent colors of the oxynitrides became orange, red, dark red, or brown based on their light absorption edge. As discussed previously, the VBM positions of perovskite oxynitrides are composed of hybridized N 2p and O 2p atomic orbitals. The CB minimum (CBM) of the oxynitrides mainly consists of empty Ti 3d, Nb 4d, or Ta 5d orbitals for the

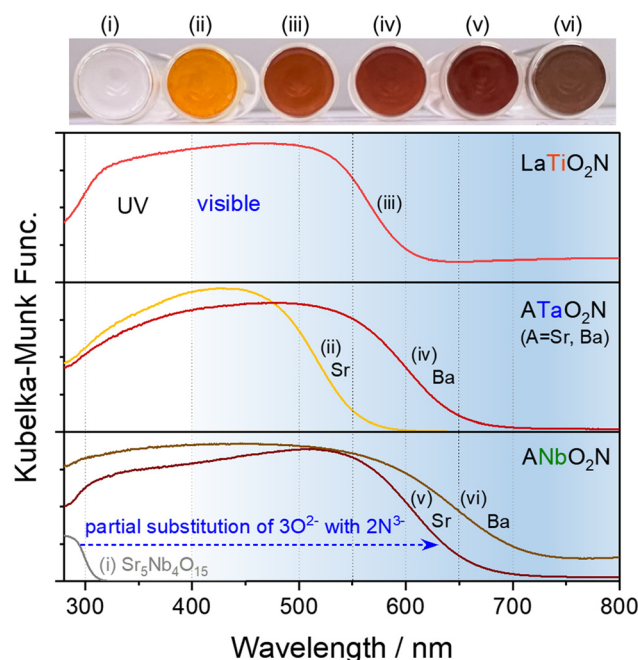


Fig. 2 Photographic images and UV-vis diffuse reflectance spectroscopy (DRS) spectra of perovskite oxynitrides $\text{AB}(\text{O},\text{N})_3$ ($\text{A} = \text{Sr}, \text{Ba}$, or La and $\text{B} = \text{Ti}, \text{Ta}$, or Nb) prepared via nitridation of layered perovskites $\text{A}_5\text{B}_4\text{O}_{15}$.

respective B-site cation with d^0 electronic configuration.^{20,21} The E_g value of oxynitrides typically decreases as the effective electronegativity of the B-site cations increases.²¹ The effective electronegativity of Nb^{5+} is larger than that of Ta^{5+} , so Nb-based oxynitrides can absorb longer wavelengths of visible light (Fig. 2). Furthermore, the CBM positions of the oxynitrides are determined by the B–O/N–B bond angles.^{21,22} The tilt of the $\text{B}(\text{O},\text{N})_6$ octahedra in a perovskite structure is larger for a smaller A-site cation, resulting in a narrower CB. The reduced dispersion of the CBs shifts the CBM positions to a more negative potential, thereby causing an increase in E_g . Thus, the E_g values of BaBO_2N , SrBO_2N , and CaBO_2N are sequentially larger according to the size of the A-site cation ($r_{\text{Ba}} > r_{\text{Sr}} > r_{\text{Ca}}$).

In addition to the narrow bandgap, the band positions of the perovskite and layered perovskite oxynitrides straddle the water redox potential (Fig. 3). This indicates that perovskite oxynitrides acting as a single absorber thermodynamically drive overall water splitting to produce H_2 and O_2 without the need to supply additional energy for the reaction. The favorable optical properties of the oxynitrides are distinguishable from those of other n-type oxide semiconductors exclusively driving water oxidation, such as visible-light-responsive BiVO_4 ($\lambda < 530$ nm) and $\alpha\text{-Fe}_2\text{O}_3$ ($\lambda < 600$ nm).^{23–25} Moreover, since the CBM potentials of the oxynitrides are more negative than the reduction potential of CO_2 , HCOOH , carbon monoxide (CO), and methanol (CH_3OH) can be produced *via* photocatalysis. N_2 fixation to synthesize NH_3 gas is theoretically possible *via* photocatalysis with oxynitrides. Finally, since the VBM potentials of the oxynitrides are located below the potential of water oxidation reactions, they can catalyze the oxidation of various anions in seawater (including mineral salts).

Over several decades, perovskite and layered perovskite oxynitrides have been regarded as promising visible-light-responsive semiconductors for efficient artificial photosynthesis to produce value-added eco-friendly and renewable energy resources such as H_2 , HCOOH , CH_3OH , and NH_3 .^{2,3,26–28} This is because the oxynitrides are chemically stable in aqueous

electrolytes at various pH values and have ideal optical properties due to a narrow bandgap and a suitable band position, which makes them favorable for sunlight-driven photocatalysis. Therefore, we discuss recent advances in photocatalysis using various perovskite and layered perovskite oxynitrides responsive to a wide wavelength range of the solar spectrum. How the photoactivity of these compounds can be enhanced by their synthesis method for use in photoelectrodes, bulk and surface engineering by doping (or substitution), controlling their morphology, and crystal facet engineering is discussed in detail. We also discuss the challenges associated with using perovskite and layered perovskite oxynitrides for efficient artificial photosynthesis.

2. Visible-light-driven photocatalysis

A representative photocatalytic process using perovskite or layered perovskite oxynitrides is water splitting to produce renewable H_2 energy. The half water oxidation of water splitting typically exhibits very slow kinetics compared with half water reduction so that alternative oxidations using the oxynitrides have been addressed to boost water splitting and then produce H_2 gas with a higher evolution rate.²⁹ Recently, CO_2 reduction using these compounds has also been reported since carbon neutrality is becoming critical globally.^{30,31} First, we discuss the various reactions that can be photocatalyzed by perovskite and layered perovskite oxynitrides under sunlight (or only visible-light irradiation), the standard reduction potentials for which are reported in Table 1.

2.1. Overall water splitting *via* one-step photoexcitation

The most ideal means of producing H_2 is overall water splitting using a single semiconductor capable of absorbing visible light. Water redox reactions, *i.e.*, hydrogen evolution reaction (HER) and oxygen evolution reaction (OER), are simultaneously activated at the separate surfaces of a semiconductor particle *via*

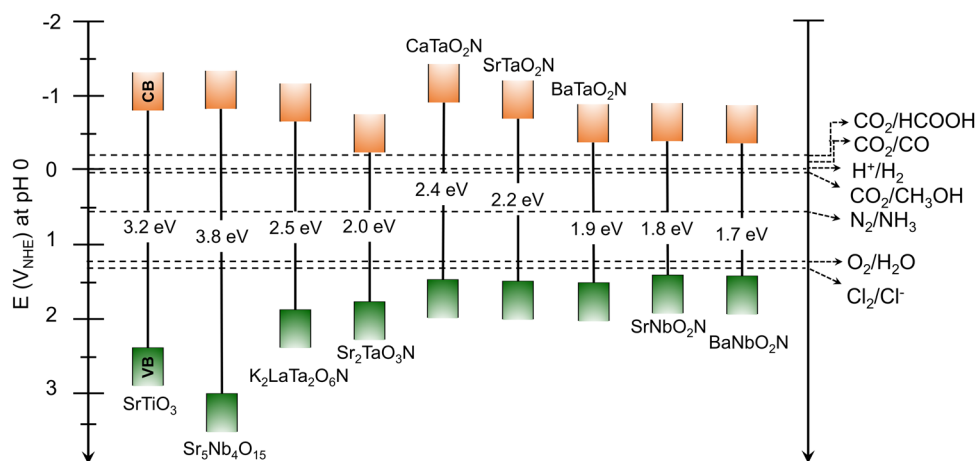
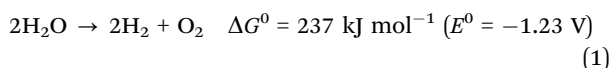


Fig. 3 Bandgap energies (eV) and band edge potentials (V_{NHE}) of representative n-type perovskite and layered perovskite oxynitrides in contact with aqueous electrolytes at pH 0. The electrochemical potentials for H_2 and O_2 evolution, CO_2 reduction, Cl_2 evolution, and N_2 fixation are also presented for comparison.

Table 1 The thermodynamic redox potentials of various reactions that can be photocatalyzed by perovskite and layered perovskite oxynitrides in contact with aqueous electrolytes at pH 0 under sunlight

Half redox reactions	Balanced chemical reactions	Potential (V_{NHE})
CO_2 reduction	$\text{CO}_2 + \text{e}^- \rightleftharpoons \text{CO}_2^-$	-1.90
	$\text{CO}_2 + 2\text{H}^+ + 2\text{e}^- \rightleftharpoons \text{HCOOH}$	-0.20
	$\text{CO}_2 + 2\text{H}^+ + 2\text{e}^- \rightleftharpoons \text{CO} + \text{H}_2\text{O}$	-0.11
	$\text{CO}_2 + 4\text{H}^+ + 4\text{e}^- \rightleftharpoons \text{HCHO} + \text{H}_2\text{O}$	-0.07
	$\text{CO}_2 + 6\text{H}^+ + 6\text{e}^- \rightleftharpoons \text{CH}_3\text{OH} + \text{H}_2\text{O}$	0.03
	$2\text{CO}_2 + 12\text{H}^+ + 12\text{e}^- \rightleftharpoons \text{C}_2\text{H}_5\text{OH} + 3\text{H}_2\text{O}$	0.08
	$\text{CO}_2 + 8\text{H}^+ + 8\text{e}^- \rightleftharpoons \text{CH}_4 + 2\text{H}_2\text{O}$	0.17
	$2\text{H}^+ + 2\text{e}^- \rightleftharpoons \text{H}_2$	0
Water reduction (HER)	$\text{N}_2 + \text{H}^+ + \text{e}^- \rightleftharpoons \text{N}_2\text{H}$	-3.2
N_2 reduction	$\text{N}_2 + 2\text{H}^+ + 2\text{e}^- \rightleftharpoons \text{N}_2\text{H}_2$	-1.1
	$\text{N}_2 + 4\text{H}^+ + 4\text{e}^- \rightleftharpoons \text{N}_2\text{H}_4$	-0.36
	$\text{N}_2 + 8\text{H}^+ + 6\text{e}^- \rightleftharpoons 2\text{NH}_4^+$	0.27
	$\text{N}_2 + 6\text{H}^+ + 6\text{e}^- \rightleftharpoons 2\text{NH}_3$	0.55
	$\text{O}_2 + 4\text{H}^+ + 4\text{e}^- \rightleftharpoons 2\text{H}_2\text{O}$	1.23
Water oxidation (OER)	$\text{Cl}_2 + 2\text{e}^- \rightleftharpoons 2\text{Cl}^-$	1.36
Chloride oxidation (CER)	$\text{ClO}_4^- + 8\text{H}^+ + 8\text{e}^- \rightleftharpoons \text{Cl}^- + 4\text{H}_2\text{O}$	1.39
	$\text{ClO}_3^- + 6\text{H}^+ + 6\text{e}^- \rightleftharpoons \text{Cl}^- + 3\text{H}_2\text{O}$	1.45
	$\text{HClO} + \text{H}^+ + 2\text{e}^- \rightleftharpoons \text{Cl}^- + \text{H}_2\text{O}$	1.48
	$\text{HClO}_2 + 3\text{H}^+ + 4\text{e}^- \rightleftharpoons \text{Cl}^- + 2\text{H}_2\text{O}$	1.57
	$\text{H}_2\text{O}_2 + 2\text{H}^+ + 2\text{e}^- \rightleftharpoons 2\text{H}_2\text{O}$	1.78
Hydrogen peroxide production	$\text{S}_2\text{O}_8^{2-} + 2\text{e}^- \rightleftharpoons 2\text{SO}_4^{2-}$	2.01
Sulfate oxidation		

one-step photoexcitation to produce H_2 and O_2 , respectively, as follows:



where E^0 is the thermodynamic potential difference for overall water splitting and ΔG^0 is the Gibbs free energy change. Thus, a semiconductor particle to drive water splitting must have an E_g greater than 1.23 eV and its band position must span the water redox potentials of 0 and 1.23 V_{NHE} . As shown in Fig. 3, the thermodynamic characteristics of almost all perovskite and layered perovskite oxynitrides satisfy these requirements, thus making them promising semiconductor candidates for water splitting *via* one-step photoexcitation.

The solar-to-hydrogen (STH) conversion efficiency (η), a benchmark figure used to estimate the performance of a semiconductor for solar water splitting, is defined by

$$\frac{\text{output energy as H}_2}{\text{energy of incident solar light}} = \frac{r_{\text{H}_2} (\text{mmol H}_2 \text{ s}^{-1}) \times \Delta G^0 (237\,000 \text{ J mol}^{-1})}{P_{\text{sun}} \times S (\text{cm}^2)} \quad (2)$$

where r_{H_2} is the rate of hydrogen evolution, P_{sun} is the photon flux of sunlight, and S is the photoreaction area of a panel or electrode. The STH efficiency greatly depends on the E_g of the semiconductor, because E_g determines the photon flux available for water splitting. Fig. 4 presents the relationships between the E_g values of perovskite oxynitrides and their theoretical maximum STH efficiency, photocurrent, and resulting hydrogen evolution rate per unit area (cm^2). Visible light in the wavelength range of 400–800 nm provides a large proportion of solar energy, and so the STH efficiency is higher in semiconductors with a wider light absorption edge. It is also known that an STH efficiency of 10% and above is required for

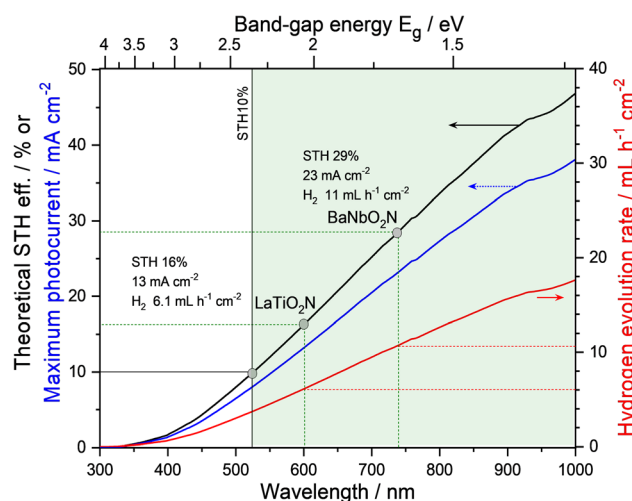


Fig. 4 A graphical representation of the theoretical maximum solar-to-hydrogen (STH) conversion efficiency, maximum photocurrent density, and hydrogen evolution rate as a function of wavelength. Representative perovskite oxynitrides are positioned along the theoretical STH efficiency curve at their respective absorption edges. The three numerical values beside each perovskite oxynitride indicate the efficiency, photocurrent density, and hydrogen evolution rate.

practical water-splitting applications.^{32,33} To satisfy this requirement, the semiconductor as a single absorber should be capable of absorbing visible light up to *ca.* 520 nm without performance loss of the water-splitting device. It is thus noteworthy that besides their suitable band potentials, perovskite and layered perovskite oxynitrides have a small enough E_g to achieve a high STH efficiency for water splitting. For instance, perovskite LaTiO_2N ($E_g = 2.1 \text{ eV}$) can adsorb visible light up to a wavelength of approximately 600 nm (equivalent to a theoretical STH efficiency of 16%). Moreover, LaTiO_2N on its own or in a

water-splitting device produces hydrogen gas at $6.1 \text{ mL h}^{-1} \text{ cm}^{-2}$, with a maximum photocurrent density of 13 mA cm^{-2} . The total photon energy harvested by BaNbO_2N (with an absorption edge of approximately 740 nm) for solar water splitting is much greater than that by LaTiO_2N , resulting in a maximum STH efficiency of 29% corresponding to hydrogen production of $11 \text{ mL h}^{-1} \text{ cm}^{-2}$. Thus, it is necessary to develop perovskite and layered perovskite oxynitrides with long light absorption edges of above 600 nm to realize commercial water-splitting applications with a high STH value.

Overall water splitting with a quantum efficiency of almost unity using one-step photoexcitation of cuboidal Al-doped SrTiO_3 ($\text{SrTiO}_3\text{:Al}$) particles with a perovskite crystal structure has been reported.¹² The remarkable water-splitting activity was attributed to the selective depositions of $\text{Rh/Cr}_2\text{O}_3$ (for HER) and a CoOOH cocatalyst (for OER) at the (100) and (110) crystal facets of $\text{SrTiO}_3\text{:Al}$, respectively. HER and OER could be separated and strongly promoted by the anisotropic transport of photogenerated electrons and holes in the single crystal. Overall water splitting on a 100 m^2 panel comprising modified $\text{SrTiO}_3\text{:Al}$ particles is an example of a safe mass hydrogen production system with a maximum STH of 0.76%.³⁴ Although the system design of the panel for scale-up is simple and cost-effective, the low STH efficiency for mass water splitting is still a significant limitation.

Nevertheless, several semiconductors capable of absorbing visible light, such as graphitic- C_3N_4 ($E_g = 2.8 \text{ eV}$),³⁷ $(\text{Ga}_{1-x}\text{Zn}_x)\text{-(N}_{1-x}\text{O}_x)$ ($E_g = 2.4 \text{ eV}$),³⁸ Ta_3N_5 ($E_g = 2.1 \text{ eV}$),³⁹ and $\text{Y}_2\text{Ti}_2\text{O}_5\text{S}_2$

($E_g = 1.9 \text{ eV}$),⁴⁰ have been successfully applied to overall water splitting *via* one-step photoexcitation. Moreover, several perovskite oxynitrides have also been utilized. Complex $\text{LaB}(\text{O},\text{N})_3$ ($\text{B} = \text{Ta}$ or Nb) solid solutions were successfully employed in the overall water splitting, resulting from the precise tuning of both E_g and bandgap positions of the oxynitrides *via* the substitution of B^{5+} with Mg^{2+} .^{41,42} These results are discussed in detail in the following section concerning doping (or substitution). Recently, overall water splitting under visible-light irradiation has also been achieved using ATaO_2N ($\text{A} = \text{Sr}, \text{Ba}$) particles with few defects.^{35,36} Fig. 5 represents the first demonstration of overall water splitting by particulate ATaO_2N crystals and the corresponding strategies based on the deposition of active cocatalysts. Under Xe lamp irradiation ($\lambda > 420 \text{ nm}$), $\text{CrO}_x/\text{Ru}/\text{IrO}_2/\text{SrTaO}_2\text{N}$ particles, which are more photoactive than BaTaO_2N , split water to produce H_2 and O_2 gases with initial evolution rates of 9.1 and $3.0 \mu\text{mol h}^{-1}$, respectively.³⁶ The apparent quantum yield of surface-modified SrTaO_2N at the wavelength of $420 \pm 30 \text{ nm}$ was 0.34% in the initial stage of water splitting, which became decreasingly saturated to 0.005% during the reaction over 48 h.

Although the H_2 evolution rate using ATaO_2N particles is very limited compared with that using $\text{SrTiO}_3\text{:Al}$ particles,¹² it is remarkable that overall water splitting using the perovskite oxynitrides is achieved under visible-light illumination. Interestingly, a general strategy for designing oxynitride particles was employed for overall water splitting *via* one-step photoexcitation. Both perovskite oxynitrides were initially prepared

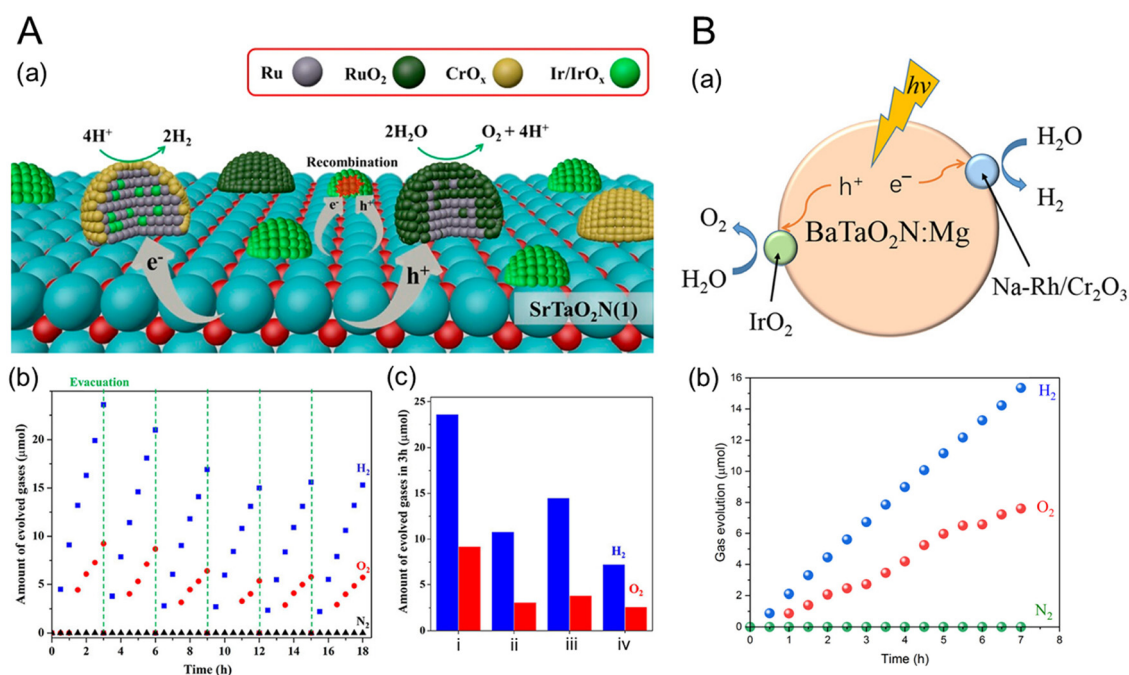


Fig. 5 Overall water splitting by (A) perovskite $\text{CrO}_x/\text{Ru}/\text{IrO}_2/\text{SrTaO}_2\text{N}$ and (B) $\text{IrO}_2/\text{Cr}_2\text{O}_3/\text{Na-Rh}/\text{BaTaO}_2\text{N:Mg}$. (a) Deposition strategies for the HER and OER cocatalysts and (b) the corresponding gas evolution activities over time under visible-light irradiation ($\lambda > 420 \text{ nm}$) using a 300 W Xe lamp in ultrapure water under an Ar atmosphere with an initial background pressure of 5 or 10 kPa. (c) The amounts of H_2 and O_2 gases generated during water splitting for 3 h using SrTaO_2N with various cocatalysts: (i) $\text{CrO}_x/\text{Ru}/\text{IrO}_2$, (ii) CrO_x/Ru , (iii) $\text{CrO}_x/\text{Ru}/\text{IrO}_{2(\text{AD})}$ (AD: by adsorption of colloidal IrO_2), and (iv) $\text{IrO}_2/\text{CrO}_x/\text{Ru}$. Reproduced with permission.³⁵ Copyright 2022, ACS. Reproduced with permission.³⁶ Copyright 2023, ACS.

with few defects *via* different synthetic routes. During nitridation, SrCl₂ and NaOH as fluxes, as well as Sr and O as sources, were introduced to obtain SrTaO₂N with few defects, while doping with Mg²⁺ with an Mg/Ta ratio of 0.1 into BaTaO₂N suppressed the Ta₃N₅ phase as an impurity. Subsequently, HER and OER cocatalysts were sequentially deposited on the perovskite oxynitride particles to promote the overall water-splitting ability. Ru or Rh covered with CrO_x or Cr₂O₃ was employed as the HER catalyst. It is well-known that chromium oxides prevent the reverse reaction (*i.e.*, the oxygen reduction reaction (ORR)) at HER sites.^{43,44} RuO₂ or IrO₂ catalysts deposited on the oxynitride surface promoted OER under light irradiation. For overall water splitting, although the bare surfaces of perovskite or layered perovskite oxides with a large bandgap (*e.g.*, SrTiO₃) can be employed as OER sites for overall water splitting,^{11,45} ATaO₂N particles with a narrow bandgap are still necessary to load the OER cocatalyst. In the solid solution case, a thin coating of amorphous oxyhydroxide on the surface of RhCrO_y/LaMg_{1/3}Ta_{2/3}O₂N suppressed N₂ evolution by self-photooxidation of the oxynitride, thereby improving overall water splitting.⁴⁶ Moreover, the deposition of HER and OER cocatalysts on ATaO₂N particles effectively suppresses the decomposition of the oxynitride and separates the reaction sites. Subsequently, the separation strategy of both types of reaction sites *via* the deposition of cocatalysts is mainly ascribed to the overall water-splitting activity of a single ATaO₂N particle.

2.2. Photoelectrochemical (PEC) water splitting

PEC water splitting using semiconductor electrodes is a potential means of producing H₂ and O₂. The HER and OER sites are divided into a p-type photocathode and an n-type photoanode, respectively. Based on this, the STH efficiency of a PEC water-splitting cell is typically higher than that of the overall water splitting using a photocatalyst.⁴⁷ Various combinations of p- and n-type electrodes have been studied for PEC water splitting without supplying electricity.^{48–50} Unbiased PEC water splitting using an inverted and planar lead halide perovskite (LHP) based on the material of a perovskite solar cell (*i.e.*, an NiFe/LHP photoanode and an NiPCoP/LHP photocathode) has recently been reported to generate a high STH conversion efficiency of 10.64% with a photocurrent density of 8.65 mA cm^{−2}.⁵⁰ A PEC cell using a perovskite BaTaO₂N photoanode combined with an La₅Ti₂Cu_{0.9}Ag_{0.1}S₅O₇ photocathode has also shown potential for the unbiased overall water splitting, albeit its activity is very low.⁵¹

Eqn (2) can be rearranged for the PEC water-splitting system to provide the STH efficiency (η) as follows:

$$\eta = \frac{|J|(\text{mA cm}^{-2}) \times \eta_F \times 1.23 \text{ V}}{P_{\text{sun}}(\text{mW cm}^{-2})} \quad (3)$$

where J , η_F , and P_{sun} are the photocurrent density obtained in a two-electrode configuration, faradaic efficiency, and the power density of sunlight (100 mW cm^{−2} for AM 1.5G), respectively. According to the definition, a PEC cell must generate high photocurrent density for water splitting without any side

reactions to obtain sufficient STH efficiency. The maximum photocurrent density of the cell is dependent on the E_g of the semiconductor, and notwithstanding, both photoelectrodes must be responsive to long wavelengths of visible light, as portrayed in Fig. 4. Moreover, the onset potentials for HER and OER at the photocathode and photoanode, respectively, are critical because the STH value of a PEC cell is estimated by using the photocurrent density obtained at the potential where the plotted linear sweep voltammetry (LSV) curves for HER and OER overlap, which is otherwise known as the operation voltage.³ Thus, a photoanode (or photocathode) capable of both absorbing long wavelengths of visible light and producing a high photocurrent density at a low potential (or high potential) is required for unassisted overall water splitting with high STH efficiency. For efficient water oxidation, the half-cell STH conversion efficiency (η) of the photoanode (a standard measure) is given by

$$\eta = \frac{|J|(\text{mA cm}^{-2}) \times \eta_F \times (1.23 - V_{\text{app}})(\text{V})}{P_{\text{sun}}(\text{mW cm}^{-2})} \quad (4)$$

where J is the anodic photocurrent density, η_F is the faradaic efficiency, V_{app} is the applied potential, and P_{sun} is 100 mW cm^{−2} for AM 1.5G. Based on these requirements, n-type perovskite and layered perovskite oxynitrides have been employed as a light absorber for the photoanode in a PEC cell to progress the OER because of their favorable optical property of being responsive to long wavelengths of visible light.^{17–19,52}

The B-site cations in perovskite AB(O,N)₃ can easily be reduced to a lower oxidation state (*e.g.*, Nb⁵⁺ to Nb⁴⁺ or Nb³⁺) during high-temperature nitridation under a reducing NH₃ atmosphere owing to their high electronegativity. The reduced B-site cations cause the generation of anion defects and impurity traces that compensate for the charge imbalance, which increases the recombination of photogenerated holes and electrons during water splitting and leads to a decrease in photoactivity. In fact, the photoreaction takes place on the surface of the oxynitride, and surface defects therein negatively influence the PEC water-splitting activity.^{18,53} The sunlight-driven PEC activities of LaTiO₂N, BaNbO₂N, and BaTaO₂N photoanodes have been significantly advanced compared to the other perovskite AB(O,N)₃ by controlling the surface defect density.^{17–19} Prior to deposition of the OER electrocatalyst, Akiyama *et al.* cleaned the surfaces of synthesized LaTiO₂N particles by using mild poly(4-styrene sulfonic acid) (PSS) (Fig. 6(A)).¹⁷ This resulted in etching of the defective surface layer of the oxynitride; although longer acid treatment increased the number of fine pores in the surface, it decreased the weight of the oxynitride layer. Afterward, the CoO_x/LaTiO₂N photoanode produced a high photocurrent density of 8.9 mA cm^{−2} at 1.23 V_{RHE} in a 1 M NaOH electrolyte at pH 13.5 under AM 1.5G irradiation, which is the highest reported so far. Moreover, its PEC water-splitting activity was more than two times higher than that of the as-prepared CoO_x/LaTiO₂N photoanode. Similarly, eliminating the surface defects in an LaTiO₂N layer *via* acid treatment with aqua regia doubled the photocatalytic HER and OER activities of the oxynitride particles.⁵⁴ Therefore, the effect

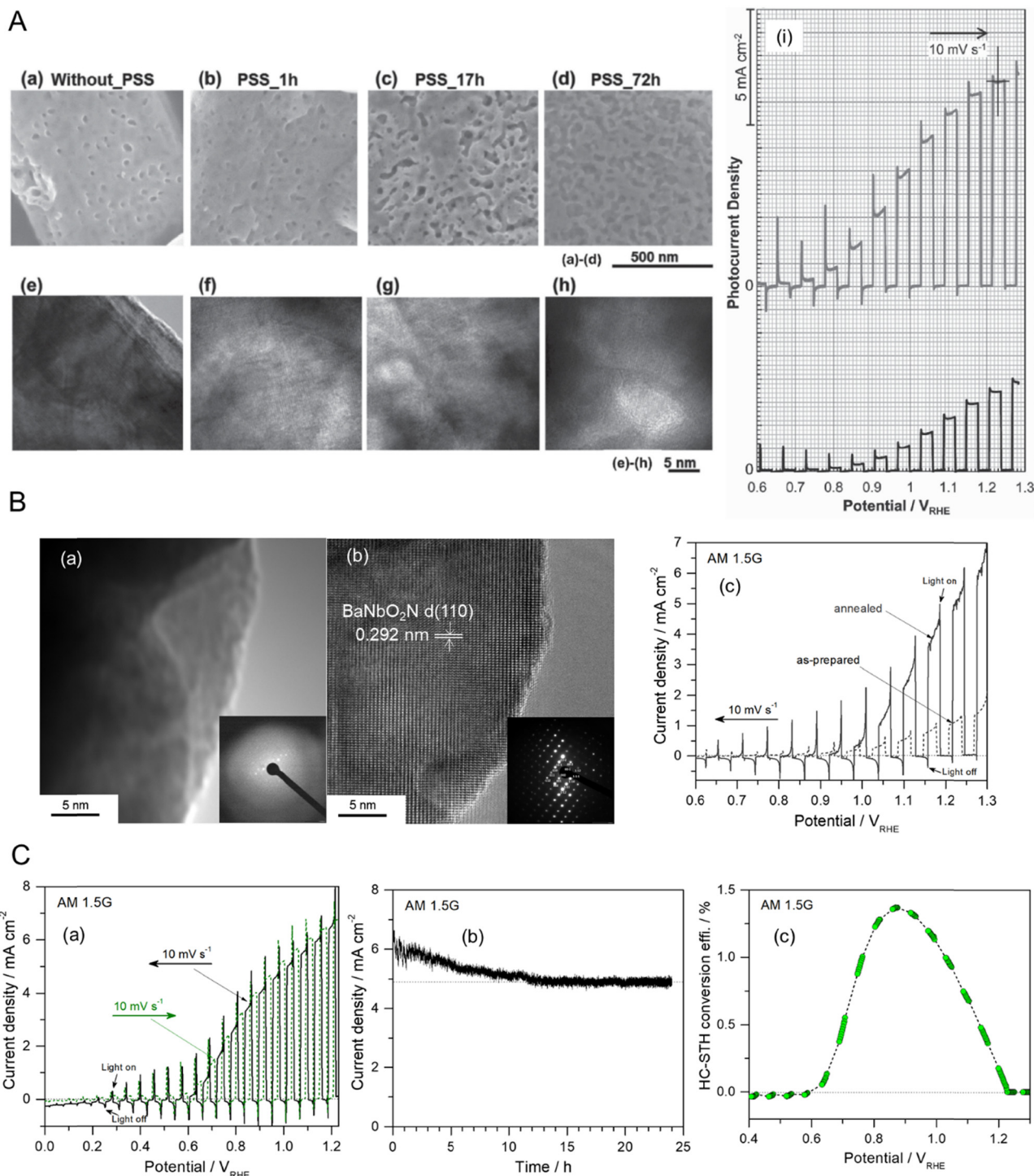


Fig. 6 PEC water splitting. (A) Scanning electron microscopy (SEM) images (a)–(d) and high-resolution (HR) transmission electron microscopy (TEM) images (e)–(h) of the surface of perovskite $LaTiO_2N$ treated with poly(4-styrene sulfonic acid) (PSS) for (a), (e) 0, (b), (f) 1, (c), (g) 17, and (d), (h) 72 h. Corresponding linear sweep voltammetry (LSV) curves of $CoO_x/LaTiO_2N$ photoanodes not treated with PSS for 17 h (upper line) used for PEC water splitting in a 1 M NaOH electrolyte at pH 13.5 under chopped AM 1.5G irradiation. Reproduced with permission.¹⁷ Copyright 2016, Wiley-VCH. (B) HRTEM images of the surface of perovskite $BaNbO_2N$ as-prepared (a) or annealed under an Ar atmosphere at 873 K for 1 h (b). Corresponding LSV curves of $Co(OH)_x-FeO_y/BaNbO_2N$ photoanodes used for PEC water splitting in a 0.5 M KBI electrolyte at pH 13 under AM 1.5G irradiation. Reproduced with permission.¹⁸ Copyright 2018, Wiley-VCH. (C) (a) LSV curves in various scan directions of a $Co(OH)_x-FeO_y/BaTaO_2N$ photoanode annealed under an Ar atmosphere at 1073 K for 1 h after nitridation used for PEC water splitting in a 0.5 M KBI electrolyte at pH 13 under AM 1.5G irradiation. (b) The chronoamperometry curve obtained during long-term water splitting at the applied potential of 1.23 V_{RHE} for 24 h. (c) The half-cell STH (HC-STH) conversion efficiency (%) estimated from the LSV curve presented in (a). Reproduced with permission.¹⁹ Copyright 2019, ACS.

of the acid treatment on photoactivity suggests that an oxynitride surface with a low number of defects is necessary to suppress the recombination of charge carriers and thereby boost water-splitting activity.

In another approach, annealing the surface of perovskite oxynitrides was effective in improving the water-splitting activity of $\text{Co}(\text{OH})_x\text{-FeO}_y/\text{BaBO}_2\text{N}$ ($\text{B} = \text{Nb, Ta}$) photoanodes under sunlight (Fig. 6(B) and (C)).^{18,19} The surfaces of the as-prepared oxynitrides were amorphous as a result of mild nitridation of Lewis base Ba-rich $\text{Ba}_5\text{B}_4\text{O}_{15}$ as the starting oxide. Subsequently, their highly defective surface with a high concentration of oxygen atoms became a single crystal by annealing under an Ar atmosphere at a suitable temperature. For instance, Ar-annealing of BaNbO_2N at 873 K for 1 h resulted in a photocurrent density of 5.2 mA cm^{-2} at $1.23 V_{\text{RHE}}$ for sunlight-driven PEC water splitting in a 0.5 M KBi aqueous electrolyte at pH 13, which is five times higher than that of the as-prepared oxynitride.¹⁸ Although using inert Ar gas has provided the most favorable results, the amorphous surface of oxynitrides can also be crystallized by annealing under other gas atmospheres, such as NH_3 . The degree of improvement in photoactivity is inversely proportional to the surface defect concentration of the as-prepared oxynitride. Moreover, a highly defective oxynitride surface can become polycrystalline rather than a single crystal *via* Ar-annealing, leading to a small increase in photocurrent. The effect of the annealing temperature on the oxynitride is also dependent on the thermal stability of the oxynitride under an Ar atmosphere.⁵³ Prolonged annealing at the decomposition temperature of BaNbO_2N ($> 873 \text{ K}$) decreases the photoactivity. For instance, Ar-annealing highly enhanced the surface crystallinity of BaTaO_2N , probably because of its high thermal stability nearly up to 1200 K. Thus, the increased crystallinity of as-prepared bulk BaTaO_2N and its surface *via* Ar-annealing at 1073 K for 1 h induced a high photocurrent density of 6.5 mA cm^{-2} at $1.23 V_{\text{RHE}}$ for water splitting in a 0.5 M KBi aqueous electrolyte at pH 13.¹⁹ The photoactivity of the oxynitride corresponded to a maximum half-cell STH energy conversion efficiency of 1.4% at $0.88 V_{\text{RHE}}$, which is still the highest value yet reported using perovskites $\text{AB}(\text{O,N})_3$. Moreover, the improved crystallinity both on the surface and in the bulk of BaTaO_2N led to long-term stability during water splitting over 24 h (79% retention of the initial photocurrent). These remarkable results clearly demonstrate that modifying the surface crystallinity of $\text{AB}(\text{O,N})_3$ *via* annealing or using acid etching to improve the surface texture, both of which lower the defect density, is an excellent way of providing strong separation and fast transfer of photogenerated charges therein, thus leading to highly active and stable PEC water splitting.

2.3. Alternative oxidation reactions to the OER

OER driven by four-electron transfer is relatively sluggish compared with HER activated *via* a two-electron pathway, the former creating a bottleneck for water splitting leading to a significant overpotential. The sluggish kinetics of OER typically become much slower in neutral electrolytes regardless of the electrocatalysts and/or semiconductors used, which indirectly causes a low production rate of hydrogen. Alternatively, a high

evolution rate of hydrogen from water at neutral pH can be achieved *via* other oxidation reactions with relatively fast kinetics,^{28,55,56} such as the chlorine evolution reaction (CER). In particular, the splitting of seawater with a high concentration (*ca.* 0.55 M) of Cl^- involving the CER can be driven under sunlight. In seawater, the thermodynamic potential for the oxidation of Cl^- driven *via* two-electron transfer to produce Cl_2 is $1.36 V_{\text{NHE}}$ at pH 0 (Table 1). Fig. 7(A) represents a calculated Pourbaix diagram for an artificial chlorine system in a 0.5 M NaCl aqueous electrolyte.⁵⁷ The disproportionation of Cl^- in water to produce hypochlorous acid (HClO), which is also activated by the two-electron pathway, is possible in the pH range of approximately 3–7.5 while the dissociation of HClO producing hypochlorite (ClO^-) occurs at neutral and alkaline pH values because of its pK_a of 7.5. Although CER is thermodynamically less favorable than OER ($E^0 = 1.23 V_{\text{NHE}}$ at pH 0), the oxidation driven *via* the two-electron pathway in the former is kinetically catalyzed faster than in the latter. The CER is selectively more catalyzed than the competing OER even in strong acidic electrolytes due to its independence from the concentration of H^+ .⁵⁸ Moreover, using seawater instead of pure water is cost-effective for artificial H_2 production systems because the former is very abundant. Based on the advantages of the CER over the OER, PEC seawater splitting using WO_3 and BiVO_4 semiconductors at neutral pH has recently been reported.^{58–60}

The photoactivity of perovskite oxynitrides for OER is relatively low in neutral electrolytes compared to strong alkaline electrolytes (*e.g.*, at pH 13).^{61–63} This is because the band edge and flat band potential energies of $\text{AB}(\text{O,N})_3$ are dependent on the pH value, and their OER kinetics are very slow under neutral conditions.^{14,53} However, the limited activity of $\text{AB}(\text{O,N})_3$ at a neutral pH makes them unfavorable for commercial water-splitting applications. Recently, the PEC activity of SrNbO_2N for water splitting at a neutral pH was largely enhanced *via* the bottom-up fabrication of the photoanode and the addition of NaCl to an aqueous electrolyte.²⁹ The feasible mechanism to simultaneously drive the HER and CER using the perovskite oxynitride in artificial seawater is illustrated in Fig. 7(B). The band edge potentials of SrNbO_2N thermodynamically span the standard electrode potential for CER, as well as those for OER and HER, indicating that the holes photogenerated at the surface of the oxynitride are consumed to drive the CER and OER. Fig. 7(C) demonstrates sunlight-driven seawater splitting in a 0.5 M NaCl aqueous electrolyte at pH 6.4 using an SrNbO_2N photoanode prepared *via* bottom-up fabrication including oxidation and flux-assisted nitridation: the oxidation process was necessary for growing the crystalline Nb_2O_5 layer on an Nb substrate while the subsequent two-step nitridation process at different temperatures caused the complete conversion of crystalline Nb_2O_5 to porous cuboidal SrNbO_2N with high crystallinity and a large surface area. Consequently, the $\text{Co}(\text{OH})_x/\text{SrNbO}_2\text{N}/\text{Nb}$ photoanode exhibited an mA-level photocurrent in a neutral 0.2 M NaPi aqueous electrolyte, which is a remarkable result for water splitting using a perovskite oxynitride. Moreover, the photoactivity of the oxynitride became three times higher in artificial

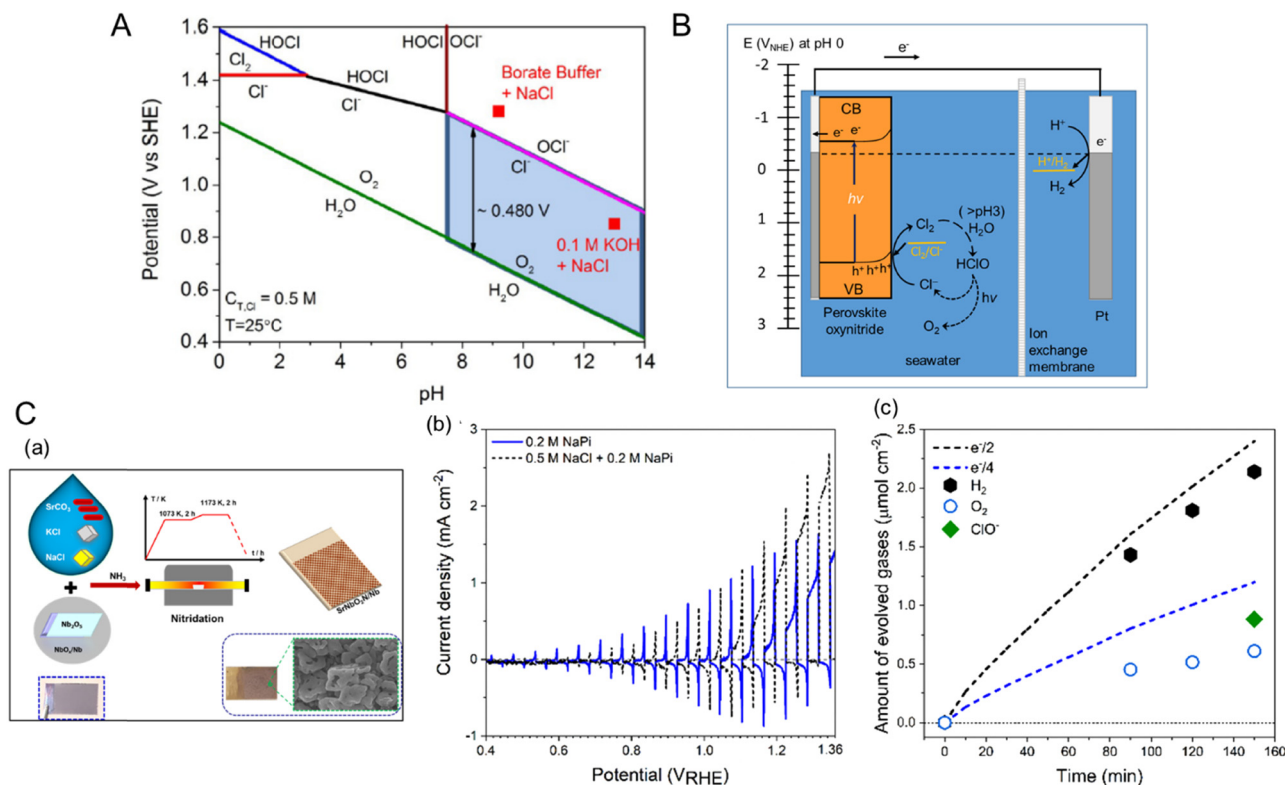
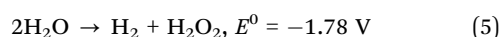


Fig. 7 Chlorine evolution reaction (CER) in seawater as an alternative oxidation pathway. (A) A computed Pourbaix diagram for an artificial chlorine system in a 0.5 M NaCl aqueous electrolyte without any other mineral salts. Reproduced with permission.⁵⁷ Copyright 2016, Wiley-VCH. (B) The feasible mechanism for PEC seawater splitting using a perovskite oxynitride in a 0.5 M NaCl aqueous electrolyte. (C) (a) A schematic of the bottom-up fabrication of an SrNbO₂N/Nb photoanode involving oxidation and two-step nitridation. (b) LSV curves for the corresponding Co(OH)_x/SrNbO₂N/Nb photoanode during seawater splitting in a 0.2 M NaPi buffer with and without 0.5 M NaCl electrolyte (at pH 6.4 and 6.7, respectively) under AM 1.5G irradiation. (c) Time courses of O₂, ClO₂⁻, and H₂ generation during seawater splitting using the same photoanode and a Pt wire at an applied potential of 1.23 V_{RHE} in a 0.5 M NaCl electrolyte buffered with 0.2 M NaPi under AM 1.5G simulated sunlight for 150 min. The dashed lines indicate the amounts of H₂, ClO₂⁻ (e⁻/2), and O₂ (e⁻/4) estimated for a faradaic efficiency of unity. Reproduced with permission.²⁹ Copyright 2023, ACS.

seawater including 0.5 M NaCl providing the activation of the CER as well as the OER. Quantitative analysis proved that seawater splitting by SrNbO₂N resulted in the generation of HClO, ClO⁻, and O₂ in the oxidation compartment and H₂ evolution in the reduction compartment, corresponding to a faradaic efficiency of almost 90%. These results clearly indicate that the CER is more preferentially driven over the OER during seawater splitting at neutral pH. It also indicates that the CER can be an alternative oxidation reaction to the OER that improves the H₂ production activity of perovskite oxynitrides in strong alkaline to neutral environments.

Hydrogen peroxide (H₂O₂) production from water oxidation is regarded as an alternative oxidation reaction to the OER,^{28,64} and H₂O₂ is also generated *via* the ORR using semiconductor photocathodes.^{65,66} In this review, we only discuss H₂O₂ production from water oxidation in terms of improving H₂ production. As reported in Table 1, although H₂O₂ production is driven *via* a two-electron pathway similar to the CER leading to kinetically fast activation, its thermodynamic potential of 1.78 V_{NHE} is significantly unfavorable compared with the competing OER. The overall reaction for H₂O₂ production and water reduction *via* one-step photoexcitation is as follows:



Despite the thermodynamic disadvantage, the production of H₂O₂ based on the photocatalytic oxidation of water has recently been attempted because H₂O₂ is a much higher value-added product than O₂.^{67–69} Moreover, attempts at suppressing the competitive OER and selectively improving the valuable H₂O₂ production have been successful. In particular, it has been reported that the HCO₃⁻ electrolyte in the pH range of 7–8 promotes the PEC activity of a BiVO₄/WO₃ photoanode for H₂O₂ production.⁷⁰ However, the very low faradaic efficiency for the PEC H₂O₂ production (approximately 54%) was improved up to 79% by introducing an Al₂O₃ overlayer on the photoanode.⁶⁷ In the reaction mechanism, the weakly basic HCO₃⁻ as a hole acceptor is adsorbed at the weakly acidic Al₂O₃ surface and oxidized to unstable HCO₄⁻, which then reacts with H₂O to produce H₂O₂. Furthermore, a faradaic efficiency of more than 90% for H₂O₂ production using the Gd-doped BiVO₄ photoanode in the potential range of 1.8–2.5 V_{RHE} has been reported.⁶⁹ The authors theorized that the Gd doping of BiVO₄ shifted the binding energy of OH⁻ on the active sites of Bi–Bi to a more optimal energy level, thereby boosting H₂O₂ production. Although the catalytic mechanism for H₂O₂ production is still

uncertain,⁶⁴ it is remarkable that the selectivity and faradaic efficiency of the H_2O_2 production can be highly improved despite its unfavorable thermodynamic characteristics compared with the OER.

Although semiconductors with a wide bandgap (e.g., TiO_2 , WO_3 , and BiVO_4) have been employed as photocatalysts or photoelectrodes for H_2O_2 production,^{28,64,65} photocatalytic oxidation to produce valuable H_2O_2 using perovskite or layered perovskite oxynitrides has not yet been reported. Based on the electromotive force of H_2O_2 production (1.78 V) being higher than that of water splitting, the band structures of perovskite oxynitrides with a negative VBM potential could make them unsuitable. However, the band structures of layered perovskite oxynitrides with relatively large E_g values possibly make them suitable for H_2O_2 production (Fig. 3). Moreover, the bandgap engineering of perovskite oxynitrides *via* substitution with alien elements could make them capable of thermodynamically driving H_2O_2 production.^{41,46} Therefore, in addition to CER, H_2O_2 production using perovskite or layered perovskite oxynitrides may be an alternative oxidation reaction to OER that efficiently produces not only valuable H_2O_2 but also H_2 .

2.4. CO_2 reduction

The photocatalytic reduction of CO_2 using a semiconductor is driven *via* a multiple-electron transfer process that generates many different products, such as formic acid (HCOOH), carbon monoxide (CO), methanol (CH_3OH), methane (CH_4), and so on, depending on the number of transferred electrons and the subsequent reaction pathway (Table 1). The catalytic mechanism is complex and has been unclear until very recently. Indeed, much research effort toward understanding it and increasing the selectivity for its high-value reaction products has been made because of the current global drive toward carbon neutrality.^{71–73} Photocatalytic and PEC systems employing perovskite or layered perovskite oxynitrides for CO_2 reduction under visible-light irradiation have also been reported.^{30,74}

The band edge potentials of perovskite and layered perovskite oxynitrides straddling the various CO_2 reduction potentials

in Fig. 3 suggest their suitability as a semiconductor for sunlight-driven CO_2 reduction. In fact, the overall reaction including the CO_2 reduction and OER in (sea)water containing dissolved CO_2 catalyzed by oxynitrides is thermodynamically feasible *via* one-step photoexcitation. However, this has still not been realized because of the small driving force of oxynitrides with a narrow bandgap for CO_2 reduction and sluggish OER kinetics driven *via* the simultaneous transfer of four electrons. Alternatively, a hybrid PEC cell constructed using a $\text{CuGaO}_2/\text{PRu-Re}$ photocathode and a CoO_x/TaON photoanode exhibited visible-light-driven CO_2 reduction to produce CO and oxidation *via* the OER to release O_2 in an aqueous electrolyte with no external bias,⁷⁵ albeit the efficiency of the system was very poor.

In the study by Yoshitomi *et al.*, although triethanolamine (TEOA) in an organic solvent instead of water was used as the hole acceptor, a hybrid perovskite CaTaO_2N coupled with a binuclear $\text{Ru}(\text{II})$ complex (RuRu') catalyst activated CO_2 reduction under visible-light illumination to produce HCOOH with high selectivity.⁷⁶ Fig. 8(A) illustrates the expected Z-scheme CO_2 reduction mechanism of the hybrid catalyst under visible light. The binuclear RuRu' complex is composed of a redox photosensitizer unit ($\text{Ru}(\text{PS})$) and a catalytic unit ($\text{Ru}(\text{Cat})$). Both CaTaO_2N and RuRu' are capable of absorbing visible light up to a wavelength of 500 nm. The authors proposed that the transfer of photoexcited electrons from the CBM potential of CaTaO_2N to $\text{Ru}(\text{PS})$ and then to $\text{Ru}(\text{Cat})$ provides the Z-scheme for CO_2 reduction. The effect of each component in the hybrid catalyst on the CO_2 reduction activity is summarized in Table 2. The combination of CaTaO_2N with RuRu' exclusively enabled the detection of formate resulting from the dissociation of HCOOH with a high selectivity of greater than 99%. The oxynitride with only $\text{Ru}(\text{Cat})$ adsorbed thereon produced less formate and accompanying byproduct H_2 , while its combination with $\text{Ru}(\text{PS})$ did not produce either product. These results indicate that both CaTaO_2N and the binuclear RuRu' complex were necessary to realize HCOOH production. Moreover, the deposition of Ag particles on the CaTaO_2N surface resulted in three times higher CO_2 reduction

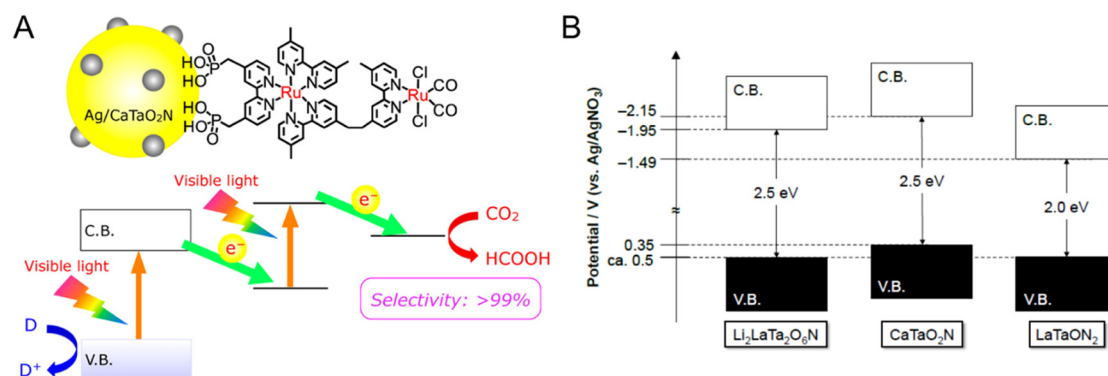


Fig. 8 CO_2 reduction. (A) The catalytic mechanism for visible-light-driven Z-scheme CO_2 reduction using a hybrid catalyst consisting of perovskite and layered perovskite oxynitrides and a binuclear Ru complex (RuRu'). Reproduced with permission.⁷⁶ Copyright 2015, ACS. (B) Band structure diagrams of layered perovskite $\text{Li}_2\text{LaTa}_2\text{O}_6\text{N}$ and perovskites CaTaO_2N and LaTaON_2 estimated in an anhydrous acetonitrile (MeCN) electrolyte containing 0.1 M tetraethylammonium tetrafluoroborate (Et_4NBF_4) at pH 7. Reproduced with permission.²⁶ Copyright 2018, Wiley-VCH.

Table 2 The photocatalytic CO₂ reduction activity of various hybrid catalysts consisting of perovskite or layered perovskite oxynitrides and a binuclear Ru complex (RuRu'). Reproduced with permission.²⁶ Copyright 2018, Wiley-VCH. Reproduced with permission.⁷⁶ Copyright 2015, ACS

Parameter	Photocatalyst ^a	Product amount (nmol)		Selectivity for formate (%)
		Formate	H ₂	
Reaction medium	CaTaO ₂ N	ND	ND	—
	Ag/CaTaO ₂ N	ND	ND	—
	RuRu'	ND	ND	—
	RuRu'/CaTaO ₂ N	93	ND	—
Ag	RuRu'/Ag/CaTaO ₂ N	320	ND	> 99
	Ru(Cat)/Ag/CaTaO ₂ N	114	2.6	—
RuRu'	Ru(PS)/Ag/CaTaO ₂ N	ND	3.8	—
	RuRu'/Li ₂ LaTa ₂ O ₆ N	ND	ND	—
Semiconductor ^b	RuRu'/Li ₂ LaTa ₂ O ₆ N	660	16	97
	RuRu'/Ag/Li ₂ LaTa ₂ O ₆ N	1440	16	99

^a Reaction conditions: CO₂ reduction for 15 h using 4.0 mg of photocatalyst in a CO₂-purged *N,N*-dimethylacetamide (DMA)/triethanolamine (TEOA) (4 : 1 v/v) electrolyte of 4 mL under a 400 W high-pressure Hg lamp with a filter comprising sodium nitrite (NaNO₂) solution. In each case, the adsorbed amount of RuRu' and the loading amount of Ag on the semiconductor were 2.5 μmol g⁻¹ and 1.0 wt%, respectively. ^b The semiconductor upon which 3 μmol g⁻¹ of RuRu' was adsorbed was immersed in mixed anhydrous acetonitrile (MeCN)/TEOA (4 : 1 v/v). RuRu' is composed of a redox photosensitized unit (Ru(PS)) and a catalytic unit (Ru(Cat)). ND, not detected.

activity. The metallic Ag particles did not act as a cocatalyst but instead as an intermediate promoter by accumulating photo-generated electrons and then mediating charge transfer from CaTaO₂N to RuRu'. This high activation and selectivity for visible-light-driven CO₂ reduction can be attributed to fast electron transfer caused by the suitable arrangement of the energy levels of CaTaO₂N and RuRu' and by the deposition of Ag particles.

It has been reported that visible-light-driven CO₂ reduction can be highly activated by introducing a 2D-layered perovskite Li₂LaTa₂O₆N instead of CaTaO₂N.²⁶ Although the *E_g* value of layered perovskite Li₂LaTa₂O₆N is the same as that of CaTaO₂N, its band structure is slightly shifted to a more positive potential from that of the perovskite oxynitride due to its different A-site cations and lower concentration of nitrogen (Fig. 8(B)). According to the CO₂ reduction activity reported in Table 2, the CBM potential of Li₂LaTa₂O₆N, even though reduced, is sufficient to transfer photogenerated electrons to RuRu'. The tendency for the photoactivity of Li₂LaTa₂O₆N is identical to that of CaTaO₂N, *i.e.*, the combination with RuRu' and the deposition of Ag nanoparticles thereon are essential for driving the photo-reaction with high selectivity. Interestingly, the CO₂ reduction activity using Li₂LaTa₂O₆N was almost five times higher than that using CaTaO₂N. The largely enhanced photoactivity was mainly ascribed to the high crystallinity of Li₂LaTa₂O₆N with a lower density of defect traps and higher density of reactive electrons, as analyzed by using transient absorption spectroscopy. However, the effect of the 2D structure of Li₂LaTa₂O₆N on the photoactivity has not yet been elucidated.^{9,77}

Perovskite oxynitrides employed as a single photocatalyst rather than in a hybrid catalyst with organic compounds for visible-light-driven CO₂ reduction have recently been reported.^{78–80} The deposition of a core-shell Ni-Ag bicomponent cocatalyst on CaTaO₂N increased interfacial electron transfer and thus enhanced the CO₂ reduction activity, showing a synergistic effect of the bicomponent catalyst.⁷⁸ Moreover, heterojunction structures composed of a metal oxide (*e.g.*, CeO₂) and LaTiO₂N have been

shown to improve CO₂ reduction, which is a well-known strategy for effectively suppressing the recombination of photo-generated charges and thereby boosting the photoreaction.^{79,80} This approach for CO₂ reduction mainly results in CO production accompanied by CH₄ as a secondary product, thereby indicating relatively low selectivity for them. Therefore, there is still room for improvement in sunlight-driven CO₂ reduction activity and product selectivity using perovskite and layered perovskite oxynitrides. Nevertheless, successful CO₂ reduction using the oxynitrides also demonstrates the suitability of their band structures for various artificial photosynthetic processes and H₂ production *via* water splitting.

3. Synthesis of perovskite and layered perovskite oxynitrides

3.1. Powder

Perovskite and layered perovskite oxynitrides are typically prepared *via* high-temperature nitridation of the starting oxides with a stoichiometric A/B ratio identical to that of the resulting oxynitride under an NH₃ atmosphere (*e.g.*, Ca₂Ta₂O₇ for CaTaO₂N, Sr₂Nb₂O₇ for SrNbO₂N, or amorphous Sr₂TaO_x for Sr₂TaO₃N).^{81–83} During nitridation, NH₃ thermally decomposes into H₂ and N₂, and N³⁻ from N₂ exchanges with O²⁻ in the starting oxide under the reducing atmosphere exemplified in the following equation for CaTaO₂N synthesis:



Otherwise, the oxynitrides are synthesized *via* calcination of the starting nitride precursor under an inert N₂/Ar atmosphere. For instance, BaTaO₂N has been prepared using TaN as the Ta precursor based on the following reaction:⁸⁴

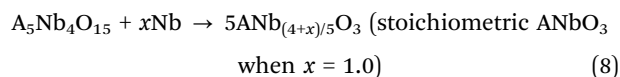


The ammonium chloride (NH₄Cl) powder instead of toxic NH₃ gas has also been used as the nitrogen source for

nitridation because it thermally decomposes into NH_3 and HCl at a temperature higher than 610 K.⁸⁵

3.1.1. Polymerized-complex method. Commonly prepared *via* a solid-state reaction (SSR), the crystalline starting oxide with a cation ratio stoichiometric to the corresponding oxynitride is completely converted to the latter according to eqn (6). For multi-metal oxynitrides such as $\text{LaMg}_x\text{Ta}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ and $\text{K}_2\text{LaTa}_2\text{O}_6\text{N}$, it is difficult to prepare the single-crystalline starting oxide *via* an SSR without traces of impurities. Fortunately, a polymerized-complex method is a facile means of preparing homogeneous starting mixed oxides at the atomic level for nitridation to oxynitrides.^{41,86,87} In short, citric acid and/or ethylene glycol are blended in methanol in which the metal precursors for the A- and B-site cations are dissolved, followed by heating to catalyze polymerization. The resulting transparent or yellow resin (proof of polymerization) is carbonized at a low temperature (≤ 773 K) to remove the organic chain in the resin. Subsequently, the carbonized powder is calcined at a higher temperature (≥ 1073 K) to finally yield a homogeneous white oxide; if it has a crystalline structure, the particle size can be controlled by varying the calcination temperature.⁸⁸ The homogeneous starting oxides are easily transformed *via* nitridation to the corresponding oxynitride with a single perovskite or layered perovskite structure that is responsive to visible light. Moreover, the resulting oxynitride is capable of water splitting and other artificial photosynthesis processes.^{26,41,42,83,89} The polymerized-complex method is effective at preparing oxynitrides with small particles and high crystallinity that provide many catalytic sites to drive photoreactions.^{41,42}

3.1.2. Conversion of A-site cation-rich starting oxides. The high crystallinity of a semiconductor is typically a critical factor for suppressing the recombination of photogenerated charges and promoting their separation and fast transfer. Many attempts to prepare perovskite and layered perovskite oxynitrides with high crystallinity and without impurity phases have been reported.^{62,91–93} It is typical that the stoichiometric oxides prepared *via* SSR form crystalline oxynitrides. If the stoichiometric oxide has a crystal structure different from the corresponding oxynitride, then the conversion of the crystal oxide phase to the perovskite or layered perovskite will definitely go ahead during nitridation. This can require a long nitridation period, which sometimes results in a low-crystalline perovskite oxynitride with a high density of defects.^{94,95} Moreover, exceptional oxynitrides such as BaBO_2N ($\text{B} = \text{Ta}, \text{Nb}$) have no corresponding starting stoichiometric crystalline oxide, so an amorphous mixture of oxides of all the elements within the oxynitride must be used instead.^{91,96} Particulate ANbO_2N ($\text{A} = \text{Sr}, \text{Ba}$) can be prepared from the nitridation of stoichiometric perovskite ANbO_3 (Nb^{4+}), as shown in Fig. 9(A).⁹⁰ Cubic perovskite SrNbO_3 is a red metallic photocatalyst for photocatalytic H_2 and O_2 evolution under visible light ($\lambda > 420$ nm).⁹⁷ Single-phase ANbO_3 can be synthesized in sealed evacuated fused quartz ampoules through the following reaction:



The SEM images in Fig. 9(B) and the XRD patterns in Fig. 9(C)-(a) show the transformation of smooth perovskite BaNbO_3 to randomly porous BaNbO_2N *via* nitridation at 1173 K for 20 h. The perovskite BaNbO_3 was monotonically converted to perovskite oxynitride BaNbO_2N without involving any structural transition, leading to the exchange of oxygen with nitrogen. Thus, the use of ANbO_3 as the starting precursor was effective in the preparation of surface- and bulk-crystalline ANbO_2N . Nevertheless, the high water-splitting activity of ANbO_2N was not achieved from stoichiometric ANbO_3 (as $x = 1.0$). The high bulk crystallinity of ANbO_2N was accompanied by an impurity phase of NbO_xN_y and a surface concentration ratio of A/Nb lower than unity, resulting in an oxynitride with rather low photoactivity. An A-site-rich oxide (as $x < 1.0$) was necessary to obtain stoichiometric oxynitrides without the impurity phase because of the volatility of the A-site cation in the presence of alkali and alkali-earth metal groups during high-temperature nitridation. In response, the use of A-site-rich precursors has been applied in various syntheses of perovskite and layered perovskite oxynitrides.^{62,94–96} Layered perovskite $\text{A}_5\text{B}_4\text{O}_{15}$ ($\text{A} = \text{Sr}, \text{Ba}$; $\text{B} = \text{Ta}, \text{Nb}$) has been employed in the synthesis of photoactive ABO_2N , as presented in Fig. 9(A).^{15,18,19} The starting oxides have a crystal structure similar to that of ABO_2N , with the A-rich concentration being replenished during high-temperature nitridation to compensate for the volatility of the A-site cations. The layered perovskite was converted to perovskite oxynitride ABO_2N *via* the following reaction:



The smooth-layered perovskites were completely changed to their porous ABO_2N counterpart *via* a process analogous to the conversion of perovskite ANbO_3 . However, they are distinguishable in that the pores are orderly located in a layered structure, probably resulting from the decomposition of an AO layer in $\text{A}_5\text{B}_4\text{O}_{15}$ and the exchange of three O^{2-} ions in the equatorial planes of octahedral BO_6 with two N^{3-} (Fig. 9(B)).⁹⁸ As shown in Fig. 9(C)-(b), (c), the transformation of $\text{A}_5\text{Nb}_4\text{O}_{15}$ to ANbO_2N was completed with no AO impurity traces after 15 h of nitridation.¹⁵ The excess of A species, undoubtedly in the form of the amorphous phase and/or nanoparticles, was easily removed with distilled water in the washing step of ANbO_2N due to the A species being a Lewis base. The Lewis base A-rich species positively suppressed the reduction of the B-site cation, leading to ABO_2N with fewer defects and an adjusted stoichiometric A/B ratio in the oxynitride. However, it caused an amorphous surface to form on ABO_2N , which became crystalline *via* subsequent Ar-annealing treatment at a suitable temperature depending on the thermal property of ABO_2N that enhanced water-splitting photoactivity. In another conversion, employing a $(\text{Na}_{1/4}\text{Ba}_{3/4})(\text{Zn}_{1/4}\text{Ta}_{3/4})\text{O}_3$ solid solution with a perovskite structure to synthesize active BaTaO_2N maximized the evaporation of volatile Na and Zn elements during nitridation.⁹⁹

3.1.3. Flux-assisted calcination/nitridation. Flux-assisted nitridation using additive molten salts, as well as the adoption of suitable starting precursors, is an effective way of preparing

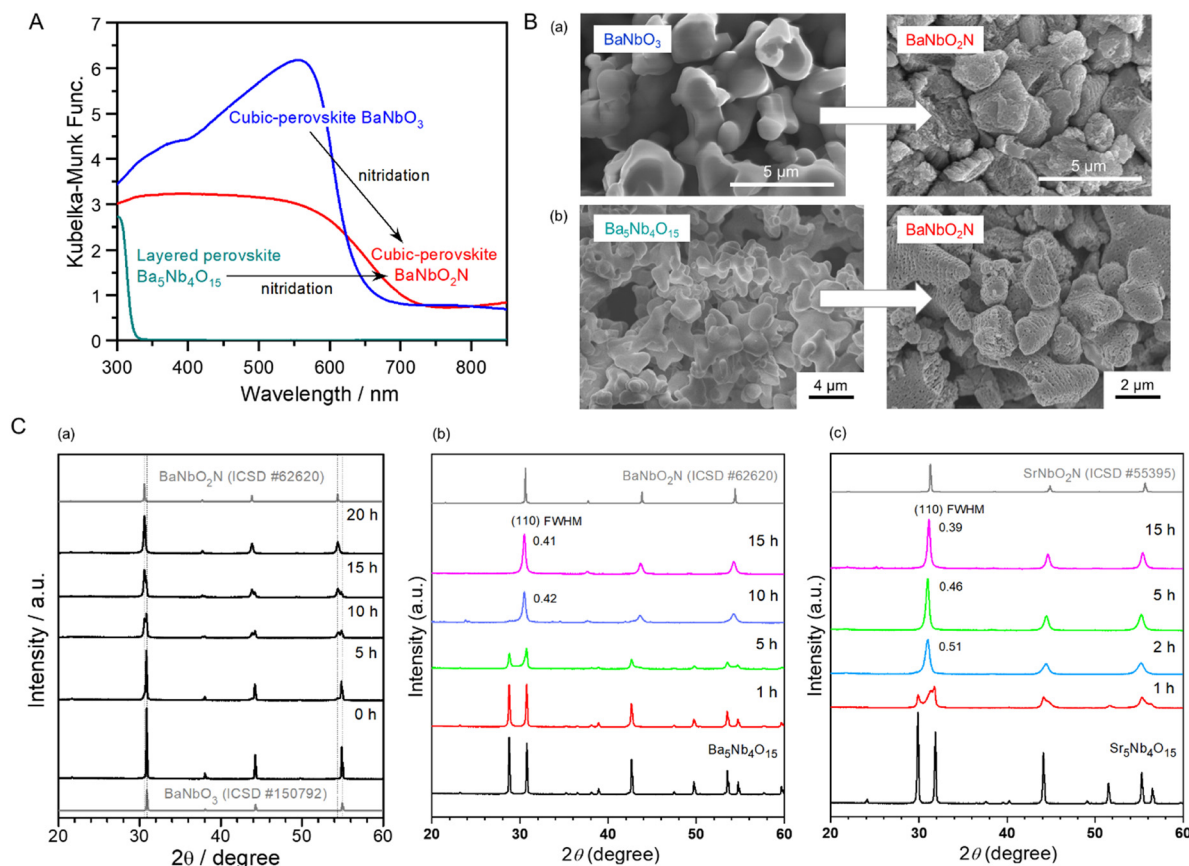
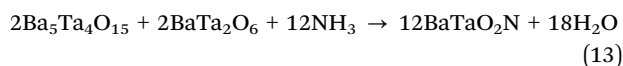
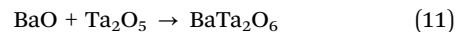


Fig. 9 Conversion of perovskite BaNbO₃ and layered perovskite Ba₅Nb₄O₁₅ to BaNbO₂N via nitridation. (A) UV-vis (DRS) spectra of starting BaNbO₃ and Ba₅Nb₄O₁₅ oxides for the synthesis of visible-light-responsive BaNbO₂N. (B) SEM images of the different starting oxides and the corresponding oxynitrides after nitridation. (C) X-ray diffraction (XRD) patterns for perovskite ANbO₂N (A = (a), (b) Ba, (c) Sr) and intermediate derivatives obtained via the nitridation of (a) BaNbO₃ and (b), (c) A₅Nb₄O₁₅ at 1123 and 1173 K, respectively. FWHM, full-width at half-maximum. Reproduced with permission.¹⁵ Copyright 2022, Elsevier. Reproduced with permission.⁹⁰ Copyright 2016, ACS.

highly crystalline perovskite and layered perovskite oxynitrides. The molten fluxes produced during high-temperature nitridation are employed as a solvent for crystallization of the oxynitrides. It is well-known that the solubility of the solute (*i.e.*, the cation precursors of oxynitrides), which determines the crystallinity and surface morphology of the resulting product (*i.e.*, the oxynitride), is influenced by the solute-to-flux molar ratio, operating temperature, and nitridation duration.^{100,101} Oxynitrides are commonly synthesized *via* the flux-assisted nitridation of the corresponding precursors in a single step or the crystalline oxide precursors prepared *via* flux-assisted calcination in two steps. Single-crystalline LaTaON₂ prepared *via* a one-step flux method using KCl exhibited higher photocatalytic and PEC activity than those prepared *via* polymerized-complex and SSR methods because of the reduced defect density and enhanced crystallinity of the oxynitride.¹⁰² Furthermore, perovskite SrNbO₂N prepared *via* one- or two-step flux-assisted nitridation became highly crystalline and the resulting oxynitride prepared from the corresponding oxide through NaI-assisted calcination provided an excellent photocurrent density of 1.5 mA cm⁻² at 1.23 V_{RHE} under AM 1.5G simulated sunlight.⁹³ Crystalline BaNbO₂N particles prepared *via* the

nitridation of Ba₅Nb₄O₁₅ and subsequent NaCl-assisted calcination provided significantly high photocatalytic OER activity under visible-light irradiation.⁹¹ These results demonstrate that flux synthesis is a worthwhile route for preparing highly crystalline perovskite oxynitrides.

Fig. 10(A) illustrates the anticipated growth mechanism of cubic-type BaTaO₂N particles showing how the flux-assisted nitridation conditions change the surface morphology of the resulting oxynitride. The following reaction steps are based on the XRD results:⁹²



First, BaCO₃ as the Ba precursor is decomposed to BaO, after which BaO and Ta₂O₅ as the Ta source are dissolved in the KCl flux at a high temperature. The diffusion of the reactants through the molten flux leads to nucleation and growth of

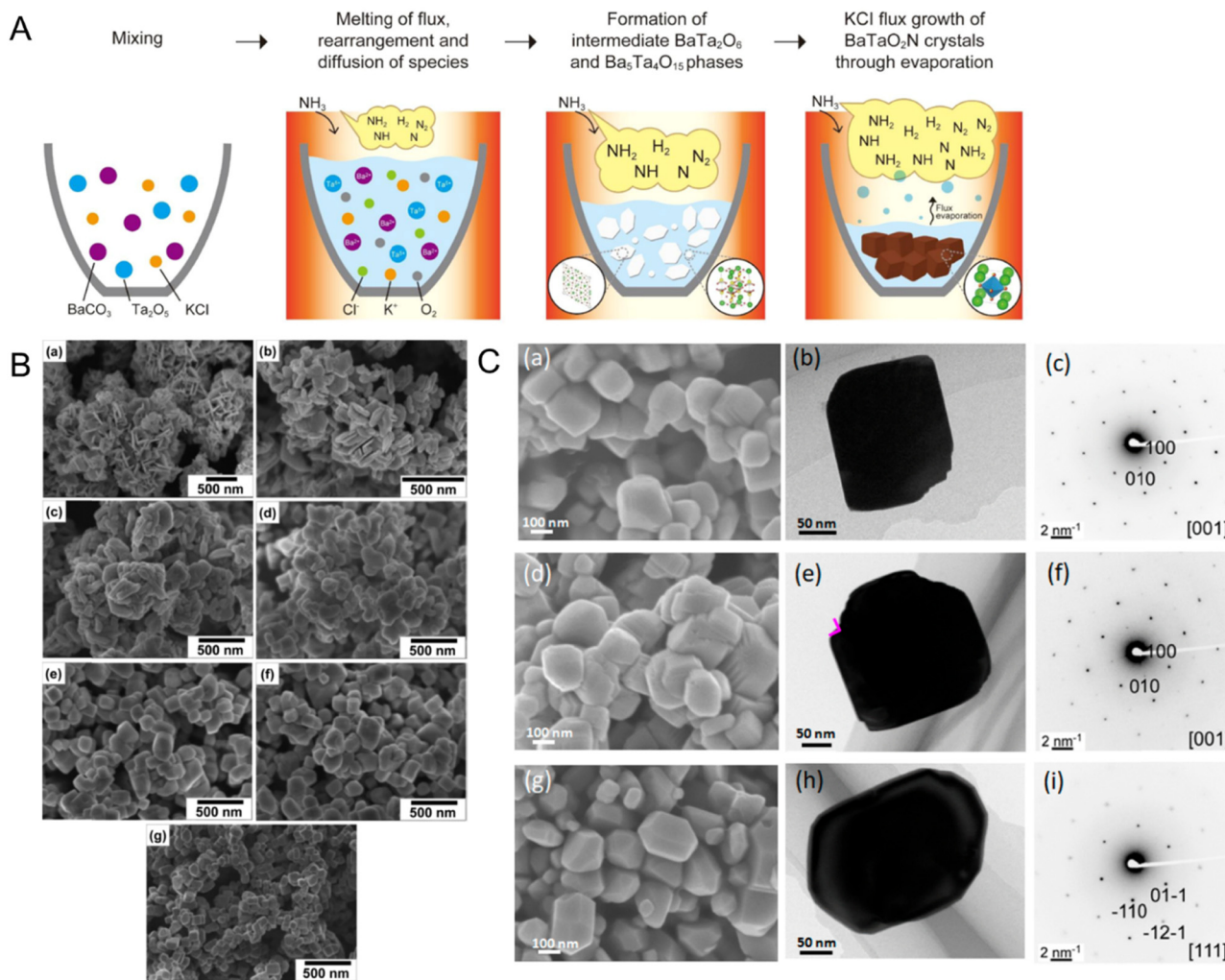


Fig. 10 Flux-assisted nitridation. (A) A schematic of the growth mechanism of cube-like BaTaO_2N crystals during nitridation using molten KCl. Reproduced with permission.⁹² Copyright 2015, ACS. (B) SEM images of BaTaO_2N particles grown *via* nitridation using molten KCl at 1223 K for (a) 0, (b) 1, (c) 2, (d) 4, (e) 6, (f) 8, or (g) 10 h. Reproduced with permission.⁹² Copyright 2015, ACS. (C) SEM images, TEM images, and SAED patterns of BaTaO_2N prepared *via* flux-assisted nitridation using different fluxes of (a)–(c) RbCl, (d)–(f) CsCl, and (g)–(i) BaCl_2 . Reproduced with permission.¹⁰³ Copyright 2020, ACS.

plate-like BaTa_2O_6 and $\text{Ba}_5\text{Ta}_4\text{O}_{15}$ (eqn (11) and (12), respectively). Finally, nitridation of the crystalline oxides under NH_3 flow causes the crystallization and growth of BaTaO_2N particles (eqn (13)). The crystalline growth during nitridation at 1123 K for 0 h (*i.e.*, as the temperature was rising) provided vertically aligned plate-like shapes (Fig. 10(B)) comprising a mixture of BaTaO_2N , $\text{Ba}_5\text{Ta}_4\text{O}_{15}$, BaTa_2O_6 , and Ta_2O_5 . The plate-like shapes became thicker and irregular with increasing nitridation time up to 4 h. After 6 h of nitridation, the irregular particles had completely turned into the intrinsic crystal shape of cubic BaTaO_2N with clear edges. Subsequently, after 10 h, the cubic-like BaTaO_2N particles displaying specific (100) and (110) facets were more homogeneously dispersed. Interestingly, the surface morphology of BaTaO_2N during nitridation with a different flux was dissimilar to that during the KCl-assisted nitridation (Fig. 10(C)).¹⁰³ As RbCl and CsCl were utilized in the latter process, BaTaO_2N particles maintained their cubic crystal

structure, although their edges were slightly truncated and the exposed facets were predominantly along the (100) plane. In particular, the BaTaO_2N crystals prepared using CeCl possessed many small steps at the edges of the cubic structure. Meanwhile, the oxynitride crystals prepared *via* BaCl_2 -assisted nitridation had a tetracaedral shape with exposed (111) and (100) facets. The various fluxes of cations changed the electrostatic forces between the precursor ions during the growth of BaTaO_2N , which thus determined the surface morphology and predominant crystal facets of the resulting oxynitride.

3.2. Photoelectrodes

3.2.1. Particulate electrodes prepared *via* spin coating, electrophoretic deposition (EPD), and particle transfer methods. Particulate powder-type perovskite and layered perovskite oxynitrides are typically synthesized *via* the nitridation of their starting precursors, as discussed previously. To prepare oxynitride

photoelectrodes for PEC artificial synthesis systems, the particulate powder must be deposited onto a conductive substrate such as F-doped tin oxide (FTO). To this end, oxynitride photoelectrodes have been fabricated *via* spin coating, EPD, and particle transfer methods.³ Spin coating is a very facile and scalable method for loading particulate oxynitrides onto conductive substrates, which has been utilized for the deposition of thin films in various applications, including solar cells.^{105–107} A suspension of the oxynitride blended with organic solvents is drop-cast on the substrate *via* a spin coater while controlling the rotation speed. The thickness of the resulting particulate film is easily controllable *via* the surface morphology of the particles, the dispersion degree of the suspension, and the rotation speed.^{106,108} For instance, 2D SrNbO_2N particles doped with Zr^{4+} have been uniformly packed onto an FTO substrate to produce a layer with a thickness of $2.5\ \mu\text{m}$ by spin coating, as shown in Fig. 11(A).¹⁰⁵ The semi-transparent oxynitride film showed a significant photocurrent density of $2.0\ \text{mA cm}^{-2}$ at $1.23\ V_{\text{RHE}}$ for water splitting under sunlight irradiation, thereby proving the feasibility of using the spin coating method to prepare water-splitting films. Meanwhile, EPD has become the most ubiquitous means of preparing photoanodes from particulate oxynitrides.^{9,95,109} Fig. 11(B) presents a schematic diagram of the fabrication of an oxynitride photoanode *via* the EPD method. The oxynitride powder is dispersed in an organic solvent including ionized species (*e.g.*, I^-), and two conductive electrodes are

immersed parallel in the suspension solution. The positively and negatively charged species are shifted to the oppositely charged electrode at an applied DC voltage *via* electrostatic attraction, and the oxynitride particles are commonly deposited on the negative electrode. The distance between the two electrodes, the concentration of the charged species, the applied voltage, the duration, and so on, can all be varied to tune the thickness and deposition density of the resulting oxynitride film. In fact, heat treatment with conductive binders to promote the transfer of photogenerated charges between multiple layers of particles and/or the substrate (called necking) is an important step in preparing photoanodes *via* EPD.^{61,95} The water-splitting photocurrent density over a TaON photoanode has been notably increased by necking treatment of the photoanode using TiCl_4 .¹¹⁰ Nevertheless, the water-splitting activity of the photoanodes prepared *via* EPD is limited due to many grain boundaries between the oxynitride particles in the multiple layers slowing the migration of photoexcited charges. It has been suggested that the particle transfer method (Fig. 11(C)) can overcome the latter disadvantage in multi-layered particulate oxynitride photoanodes.¹⁰⁴ The ohmic contact layer (typically a metal) with a thickness of a few hundred nanometers is first deposited onto the top layer of the oxynitride particles loaded on a glass substrate by using radio-frequency (RF) magnetron sputtering. A thicker Ti conductor layer ($\sim 5\ \mu\text{m}$) is then deposited on the ohmic contact layer *via* RF magnetron sputtering,

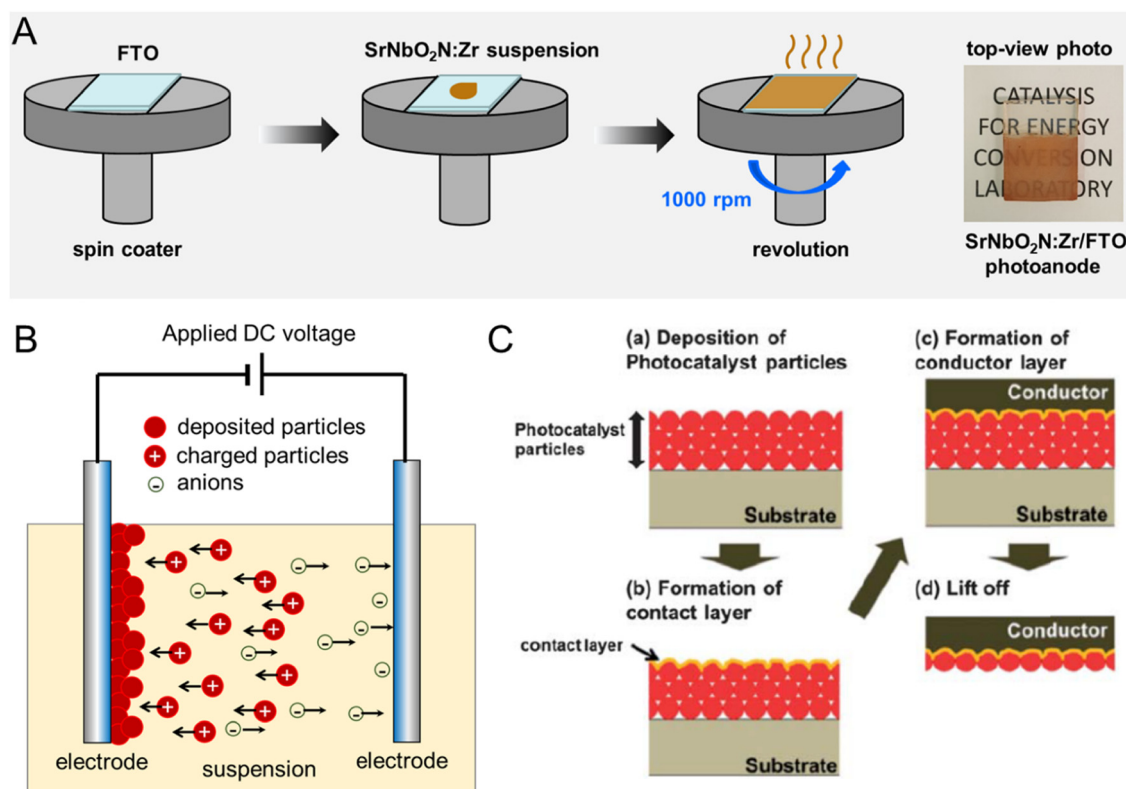


Fig. 11 Particulate photoelectrodes. Schematic diagrams of the preparation of perovskite oxynitride photoanodes *via* (A) spin coating, (B) electrophoretic deposition (EPD), and (C) particle transfer methods. Reproduced with permission.¹⁰⁴ Copyright 2013, RSC. Reproduced with permission.¹⁰⁵ Copyright 2023, Elsevier.

resulting in high electroconductivity and mechanical strength. Subsequently, several of the upper layers physically connected to the conductor and contact layers are lifted off, which finally leads to an ideal oxynitride monolayer photoanode. An $\text{LaTiO}_2\text{N}/\text{Ta}/\text{Ti}$ photoanode composed of a Ta ohmic contact and Ti conductive layers produced *via* the particle transfer method showed water-splitting photoactivity much higher than that of its counterpart prepared *via* EPD.¹⁰⁴ Moreover, the water-splitting photocurrent density at 1.2 V_{RHE} of the $\text{BaTaO}_2\text{N}/\text{Ta}/\text{Ti}$ photoanode was 10 times higher than that of previously reported photoanodes of the same material.^{61,62} The oxynitride particles on the photoanodes prepared *via* the particle transfer method are well dispersed in a monolayer, which promotes the efficient separation and vertical transfer of charges photo-generated using a single oxynitride. In fact, the highest PEC activities of perovskite oxynitrides LaTiO_2N , BaTaO_2N , and BaNbO_2N were observed in monolayered photoanode systems prepared *via* the particle transfer method.^{17–19} These findings indicate that the monolayer electrode structure resulting from the particle transfer method is effective at controlling charge separation and transfer, thus enhancing the photoactivity for water splitting.

3.2.2. Bottom-up fabrication. The bottom-up fabrication of perovskite or layered perovskite oxynitride photoelectrodes from conductive substrates increases the conductivity of photo-generated charges between the oxynitride particles and/or the substrate over that of an electrode fabricated using the oxynitride powder, which probably enhances the photoreaction capability. Based on this, various bottom-up fabrication methodologies have been introduced for preparing oxynitride photoelectrodes. LaTiO_xN_y thin films grown on a TiN conductive layer have been synthesized by using a modified pulsed laser deposition method;¹¹¹ determining the surface crystal orientation of

the (001), (011), or (112) facets of the LaTiO_xN_y thin films was achieved by using different facet-oriented TiN conductive layers. However, the water-splitting activity of the resulting oxynitride film was very limited, mainly due to the non-stoichiometric composition of the anions in the oxynitride. SrNbO_2N crystals have been grown on an Nb substrate *via* flux coating $\text{Sr}(\text{NO}_3)_2$ and chloride salts followed by nitridation.¹¹² The Sr-to-flux molar ratios and nitridation conditions determined the surface morphology and bulk crystallinity of the resulting SrNbO_2N and increased interfacial adhesion between SrNbO_2N and the Nb substrate. Despite the high interfacial adhesion of the oxynitride film, the water-splitting photoactivity was low. This was because the nitrogen during high-temperature nitridation managed to diffuse into the conductive substrate through the oxynitride semiconductor layer, which caused the generation of additional crystal and/or metallic interlayers. Thus, preparing photoactive oxynitride photoelectrodes *via* bottom-up fabrication is not straightforward.

Several studies showing the improved PEC water-splitting activity using perovskite oxynitrides prepared *via* bottom-up fabrication have recently been reported.^{52,113,114} Fig. 12 presents top-view and cross-sectional SEM images of various bottom-up-fabricated oxynitride films. Vertical SrNbO_2N nanorod arrays have been grown on an Nb substrate *via* a hydrothermal method and subsequent nitridation.⁵² SrNbO_x nanorod arrays were grown on the Nb substrate at a low temperature of 473 K, after which the oxide was transformed to SrNbO_2N nanorod arrays through nitridation at 1273 K for 2 h. As shown in Fig. 12(A), the diameter and length of the oxynitride nanorods were approximately 60 and 500 nm, respectively, thereby allowing favorable visible-light absorption and vertical separation of photogenerated charges. Interestingly, perovskite oxynitrides with 1D surface morphology have

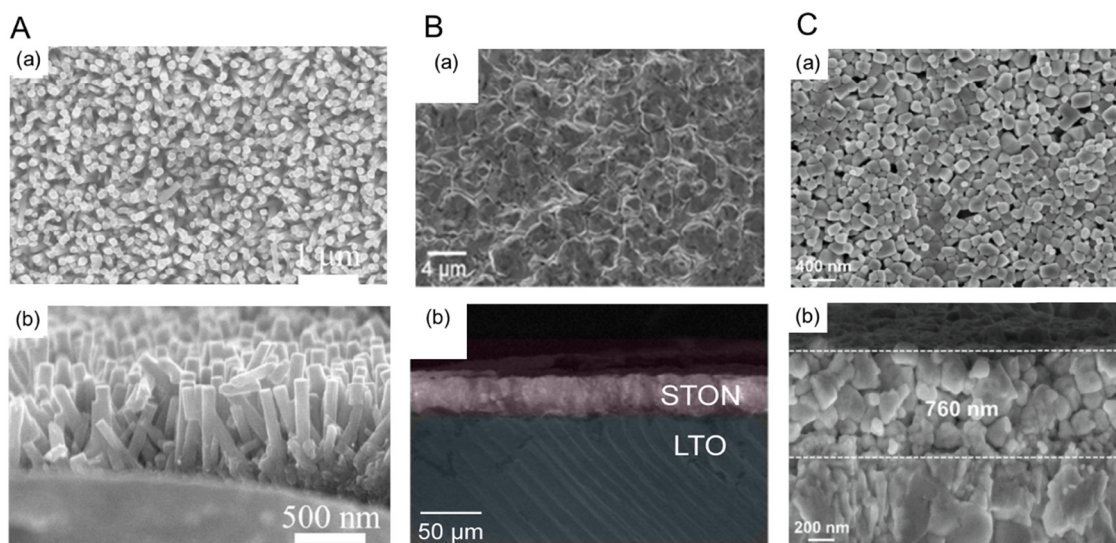


Fig. 12 Bottom-up fabrication of perovskite oxynitride photoanodes. (a) Top-view and (b) cross-sectional SEM images of (A) SrNbO_2N nanorod arrays prepared *via* a hydrothermal method followed by nitridation, (B) SrTaO_2N crystals grown on a LiTaO_3 substrate synthesized by using a reactive inorganic vapor method including nitridation, and (C) BaTaO_2N nanoparticle films deposited on a Nb substrate prepared *via* co-evaporation followed by nitridation. Reproduced with permission.⁵² Copyright 2020, Wiley-VCH. Reproduced with permission.^{113,114} Copyright 2022, Elsevier.

been reported for the first time. After the loading of CoO_x nanoparticles, SrNbO_2N nanorod arrays produced a notable photocurrent density of 1.3 mA cm^{-2} at $1.23 V_{\text{RHE}}$ in an aqueous electrolyte at pH 13 under AM 1.5G simulated sunlight. Although the synthesis of the starting oxide at low temperature and a short nitridation period caused no additional interlayers detrimental to photoreactions, the light absorption edge of the SrNbO_2N nanorod arrays was approximately 630 nm, which is significantly shorter than 700 nm reported previously.^{15,93}

Centimeter-scale perovskite SrTaO_2N crystals have been prepared by using a reactive inorganic vapor method including nitridation.¹¹⁴ The oxynitride crystals were epitaxially grown on a (110)-oriented LiTaO_3 substrate *via* the evaporation of Sr sources under an NH_3 atmosphere. During the evaporation step, Sr^{2+} was exchanged with Li^+ in the (110)-oriented LiTaO_3 and nitrogen diffused into the oxide substrate simultaneously, leading to the growth of the SrTaO_2N crystals. Interestingly, different crystal orientations of LiTaO_3 (*i.e.*, the (110) and (001) facets) resulted in nitridation rather than the intercalation of Sr^{2+} because the sizes of the lattice voids in the (110) and (001) facets are not large enough for the insertion of Sr^{2+} in the oxide substrate whereas they are in the (100) orientation. As shown in Fig. 12(B), the epitaxial growth of the perovskite SrTaO_2N crystals produced a dense film several tens of micrometers thick because of lattice expansion resulting from the insertion of Sr^{2+} . The dense film layer minimized inter-particle interfaces and grain boundaries, thereby resulting in a high photocurrent density of 1.20 mA cm^{-2} at $0.6 V_{\text{RHE}}$ (with a low onset potential of $0.35 V_{\text{RHE}}$) for water splitting in a 1 M NaOH electrolyte at pH 13.6. However, lattice expansion during the conversion to oxynitride caused the limited diffusion of Sr^{2+} , so the oxide substrate was not completely transformed to SrNbO_2N . Thus, these results indicate that the suitable selection of both the starting substrate and its crystal orientation is critical for the epitaxial growth of perovskite oxynitrides.

A co-evaporation method followed by nitridation to prepare a BaTaO_2N nanoparticle film directly grown on an Nb substrate has been reported.¹¹³ BaF_2 and Ta_2O_5 precursors were uniformly evaporated onto the substrate *via* electron beam deposition and then completely transformed to BaTaO_2N *via* nitridation under an NH_3 atmosphere. The resulting ~ 760 nm-thick BaTaO_2N film comprised nanosized cubic crystalline particles, as shown in Fig. 12(C). The high crystallinity of the oxynitride was due to the uniform dispersion of Ba and Ta atoms in the resulting film. As a result, the BaTaO_2N photoanode produced a high photocurrent density of 4.7 mA cm^{-2} at $1.23 V_{\text{RHE}}$ for water splitting in a 1 M KOH electrolyte at pH 13.6 under sunlight irradiation. The advantage of the co-evaporation method is the ability to regulate the reduction of Ta^{5+} during high-temperature nitridation by tuning the Ba/Ta ratio, which is similar to powder syntheses. In conclusion, the various bottom-up fabrication methods for perovskite oxynitrides reviewed in this article could enhance water-splitting activity by suppressing the generation of defects and additional interlayers in the resulting oxynitride film.

4. Bulk and surface engineering

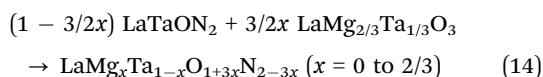
4.1. Cation doping and substitution

The doping (or substitution) of foreign elements into a semiconductor is a typical means of modifying its bulk and/or surface properties, which can lead to enhanced photocatalytic or PEC water-splitting activities. In this review, doping and substitution are distinguished from each other according to the amount of the alternative species introduced into the perovskite oxynitride: doping is below 10 at% and substitution is above 10 at%. Moreover, we cover the doping (or substitution) of the A- and/or B-site cations in the perovskite or layer perovskite oxynitrides. Perovskite BaBO_2N ($\text{B} = \text{Ta}, \text{Nb}$) has an ideal cubic crystal structure and the volume of its unit cells is markedly small.^{90,115} By contrast, the cubic lattices of $\text{AB}(\text{O},\text{N})_3$ ($\text{A} = \text{La}, \text{Ca}, \text{or Sr}$; $\text{B} = \text{Ta}, \text{Nb}$) are distorted because of the A-site cations being smaller than Ba^{2+} and the tilting of $\text{B}(\text{O},\text{N})_6$ octahedra. These findings imply that the isostructural replacement of cations in perovskite oxynitrides with alternative ones could be significantly limited in terms of the size and amount of the dopant (or substituent).

The doping of perovskite SrNbO_2N with 2 at% Zr^{4+} reduced the bulk and surface defect densities of the oxynitride by suppressing the reduction of Nb^{5+} during nitridation, thereby improving its PEC water-splitting activity.¹⁰⁵ Although the limited replacement of Nb^{5+} with Zr^{4+} maintained the optical properties of the oxynitride (*i.e.*, the absorption of visible light up to a wavelength of 680 nm), using a larger amount of Zr^{4+} caused Zr-related impurity phases, thereby notably decreasing the photoactivity. Moreover, doping the Nb^{5+} sites in BaNbO_2N with Ti^{4+} , Zr^{4+} , W^{6+} , or Mo^{6+} has been attempted to control the donor density in the oxynitride bulk originating from the generation of reduced species during nitridation.¹¹⁶ Doping with lower-valent cations Ti^{4+} and Zr^{4+} suppressed the generation of Nb^{4+} during nitridation whereas doping with higher-valent cations W^{6+} and Mo^{6+} rather increased the donor density. Moreover, doping with up to 4 at% Ti^{4+} resulted in the synthesis of BaNbO_2N without impurity traces, which thus exhibited the highest water-splitting activity. However, doping of the lower-valent cations at the Ta-sites in BaTaO_2N did not effectively lower the defect density and thus improve photoactivity despite the similarity of the ionic radii of 6-coordinate Ta^{5+} (78 pm) and 6-coordinate Nb^{5+} (78 pm). Selective replacement at the Ta-site in perovskite BaTaO_2N using 5 at% Al^{3+} , Ga^{3+} , Sc^{3+} , or Zr^{4+} for improving photocatalytic HER and OER activity has also been considered.¹¹⁷ In particular, the doping of BaTaO_2N with Mg^{2+} or Zr^{4+} enhanced OER photoactivity in a sacrificial reagent by modifying the optoelectronic and surface properties of the oxynitride. Therefore, limited doping of $\text{AB}(\text{O},\text{N})_3$ using alternative cations alters the surface properties of oxynitrides (such as defect and donor densities) while maintaining their bulk properties (such as the crystalline structure and optical properties).

Co-substitution of the Ta^{5+} site of Ta_3N_5 with 33 at% 6-coordinate Mg^{2+} (86 pm) and Zr^{4+} (86 pm) led to significant improvement in the anodic photocurrent density and onset potential for PEC water splitting.¹¹⁸ This large amount of

substitution modified not only the band structure but also the surface morphology of the original Ta_3N_5 . The difference in valence between parent Ta^{5+} and alternative cations Mg^{2+} and Zr^{4+} caused additional $\text{N}^{3-}/\text{O}^{2-}$ replacement to replenish the charge imbalance. Increasing and decreasing the concentration of $\text{Mg}^{2+}/\text{Zr}^{4+}$ cations and N^{3-} anions respectively shifted the CBM and VBM potentials negatively from those of parent Ta_3N_5 . This strategy was also applied in the synthesis of perovskite $\text{LaMg}_x\text{B}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ ($\text{B} = \text{Ta}, \text{Nb}; x = 0$ to $2/3$) solid solutions, leading to overall water-splitting activity *via* one-step photoexcitation.^{41,42} Fig. 13(A)-(a) portrays the crystal structures of $\text{LaMg}_x\text{Ta}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ solid solutions after substituting Ta^{5+} with Mg^{2+} .⁴¹ The general formula of these perovskite oxynitrides is given by



The composition of the $\text{LaMg}_x\text{Ta}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ solid solutions was easily modified by mixing perovskite oxynitride LaTaON_2 and perovskite oxide $\text{LaMg}_{2/3}\text{Ta}_{1/3}\text{O}_3$ at the atomic level. The significant substitution of Mg^{2+} in LaTaON_2 was realized after the nitridation of the starting oxides blended at the atomic level *via* a polymerized-complex method. The greater the level of substitution of Ta^{5+} with Mg^{2+} , the greater the increase in the composition

ratio of $\text{LaMg}_{2/3}\text{Ta}_{1/3}\text{O}_3$, resulting in a marked change in the optical properties of the resulting oxynitrides (Fig. 13(A)-(b)). The wavelength of the light absorption edge of $\text{LaMg}_x\text{Ta}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ blue-shifted gradually from 640 nm (pure LaTaON_2) to 270 nm (fully substituted $\text{LaMg}_{2/3}\text{Ta}_{1/3}\text{O}_3$).

This phenomenon was also observed in the substitution of Nb^{5+} with Mg^{2+} in counterpart LaNbON_2 .⁴² Fig. 13(B)-(a) presents UV-vis DRS spectra of $\text{LaMg}_x\text{Nb}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ ($x = 0$ to $2/3$) showing a gradual blue-shift of the absorption onset wavelengths from 700 to 320 nm. In addition to the increase in the E_g , the CBM and VBM potentials of the solid solutions also shifted to the negative and positive sides, respectively, as portrayed in Fig. 13(B)-(b). The band structure diagrams of the solid solution series demonstrate that the substitution of Nb^{5+} with Mg^{2+} in LaNbON_2 not only increases the concentration of the O 2p atomic orbital (leading to the positive shift of the VBM) but also decreases the concentration of the Nb 4d orbital (causing the negative shift of the CBM). The substitution of Mg^{2+} largely reduced the bulk and surface defect densities of $\text{LaMg}_x\text{Nb}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ solid solutions, which is similar to the effect of the small-cation doping of oxynitrides. Moreover, the subsequent changes in the optical properties (*i.e.*, the E_g value and band structure) of the oxynitrides were significant. In addition, surface modification using cocatalysts finally activated the overall water-splitting capability of $\text{LaMg}_x\text{B}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$,

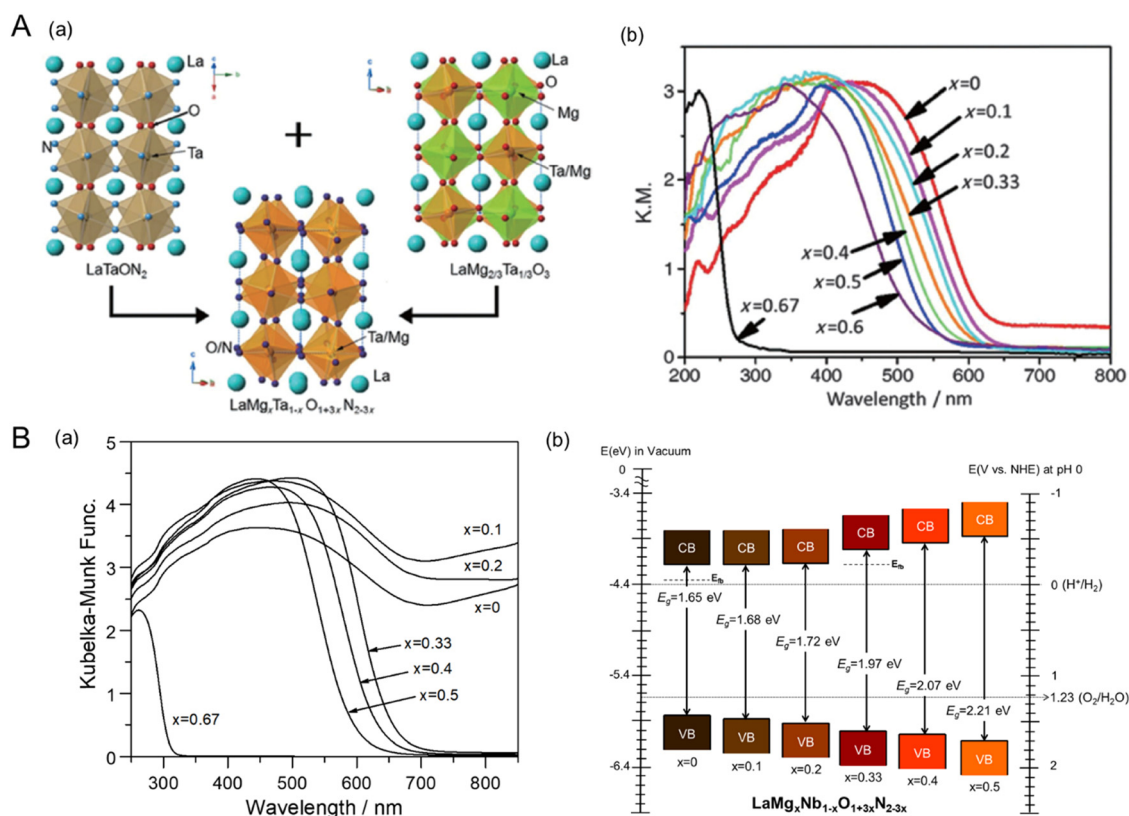


Fig. 13 Substitution of B^{5+} with Mg^{2+} in perovskite LaBON_2 ($\text{B} = \text{Ta}, \text{Nb}$). (A) (a) Crystalline structure and (b) UV-vis DRS spectra of the $\text{LaMg}_x\text{Ta}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ ($x = 0$ to $2/3$) powder after replacing some of the Ta^{5+} with Mg^{2+} . Reproduced with permission.⁴¹ Copyright 2015, Wiley-VCH. (B) (a) UV-vis DRS spectra and (b) band structure diagrams of $\text{LaMg}_x\text{Nb}_{1-x}\text{O}_{1+3x}\text{N}_{2-3x}$ ($x = 0$ to $2/3$) after replacing some of the Nb^{5+} with Mg^{2+} . Reproduced with permission.⁴² Copyright 2021, RSC.

thereby producing stoichiometric H_2 and O_2 evolution. Interestingly, the highest gas evolution rates for water-splitting were achieved using $\text{LaMg}_{1/3}\text{B}_{2/3}\text{O}_2\text{N}$ (*i.e.*, $x = 1/3$) irrespective of the cationic occupancy of the B sites in the oxynitride. As a result, the visible-light-driven overall water-splitting activity is mainly attributed to the expanded bandgap enabling a photoresponse of up to 600 and 650 nm for $\text{LaMg}_{1/3}\text{Ta}_{2/3}\text{O}_2\text{N}$ and $\text{LaMg}_{1/3}\text{Nb}_{2/3}\text{O}_2\text{N}$, respectively. Likewise, the bulk modification of oxynitrides *via* substitution with alternative cations has been subsequently applied: Zn^{2+} for Ta^{5+} in perovskite BaTaO_2N ,¹¹⁹ Zr^{4+} for Ta^{5+} in LaTaON_2 ,¹²⁰ Mg^{2+} for Ti^{4+} in LaTiO_2N ,¹²¹ La^{3+} for Ti^{4+} in layered perovskite Sr_2TiO_4 ,¹²² and so on. All these outcomes obviously demonstrate that the substitution of the B sites in oxynitrides with alternative cations is a very useful way of modifying the bulk properties of the parent oxynitride and thus improving the latter's photocatalytic water-splitting capability.

4.2. Varying the surface morphology

The electrochemical activity of an electrocatalyst (*e.g.*, Pt nanoparticles) for oxidation and reduction reactions is typically

proportional to its electrochemically active surface area containing its reaction sites for accessing the electrolyte.^{124–127} The surface morphology (including the particle size and shape) of the electrocatalyst to increase reaction sites is thus critical for progressing electrochemical reactions efficiently. In contrast, the surface morphology of a semiconductor influences visible-light harvesting, the reaction efficiency of photocarriers, and the deposition coverage of cocatalysts (*e.g.*, CoO_x) on which the actual photoreaction takes place. Because the small size of semiconductor particles can lead to a short diffusion distance for the photogenerated charges, it increases the charge separation, transfer, and collection efficiencies, thereby enhancing photoactivity.¹²⁸ BaNbO_2N particles of various sizes prepared *via* the nitridation of $\text{Ba}_5\text{Nb}_4\text{O}_{15}$ size-controlled in the range from 0.2 to 50 μm exhibit photocatalytic water oxidation activity depending on the particle size in an aqueous electrolyte including a sacrificial reagent.¹²⁹ Similarly, small scheelite-type LaNbO_4 particles as the starting oxide produced small LaNbON_2 particles with fewer defects after nitridation with significantly high photocatalytic water oxidation activity under visible-light Xe lamp

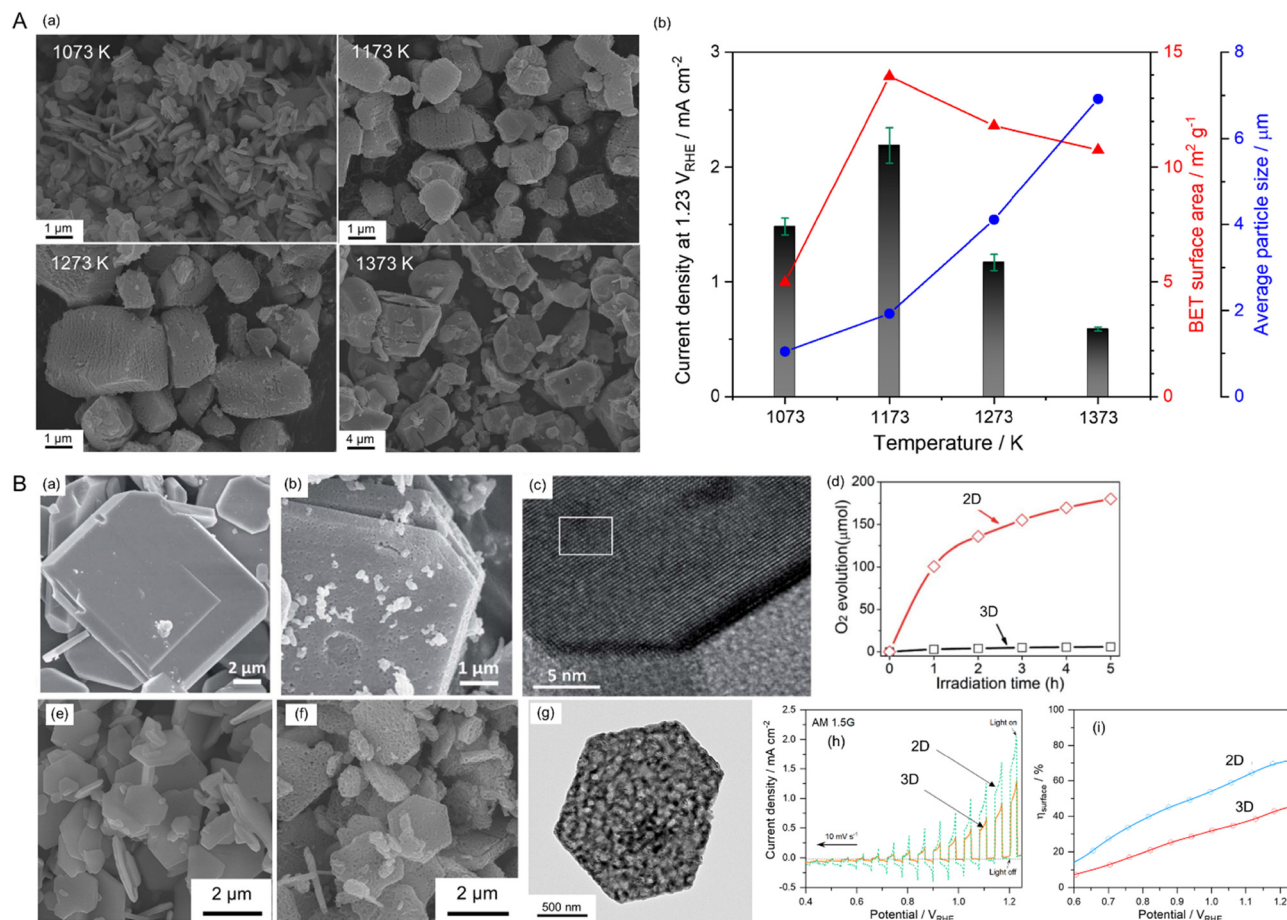


Fig. 14 Varying the surface morphology of Nb-based perovskite oxynitrides. (A) SEM images (a) and the corresponding PEC water-splitting activity (b) of BaNbO_2N particles of various sizes. Reproduced with permission.⁸⁸ Copyright 2022, Elsevier. (B) SEM images of starting LaNbON_2 (a) and the corresponding 2D LaNbON_2 (b) after nitridation along with an HRTEM image (c) and the photocatalytic O_2 evolution activity (d) of the 2D oxynitride particles. Reproduced with permission.¹²³ Copyright 2020, RSC. SEM images of starting $\text{Sr}_5\text{Nb}_4\text{O}_{15}$ (e) and the resulting 2D SrNbO_2N (f) after nitridation along with an HRTEM image (g) and the PEC water-splitting activity (h) and charge separation efficiency (i) of the 2D oxynitride photoanode. The photoactivities of both 2D oxynitrides were compared with those of typical 3D oxynitrides. Reproduced with permission.¹⁰⁵ Copyright 2023, Elsevier.

irradiation ($\lambda > 420$ nm).^{94,95} Fig. 14(A) presents the size dependence of perovskite BaNbO_2N particles on PEC water-splitting activity.⁸⁸ The size of the oxynitride particles was systematically tuned *via* the nitridation of $\text{Ba}_5\text{Nb}_4\text{O}_{15}$ with various average sizes (1.05, 1.93, 4.11, or 6.92 μm). The oxide particles synthesized *via* a polymerized-complex method were gradually grown by increasing the calcination temperature from 1073 to 1373 K. Their conversions to the corresponding oxynitride exclusively increased the surface porosity without an increase in particle size, except for the nitridation of the oxide calcined at 1073 K. The small, porous oxynitride prepared from the oxide *via* calcination at 1173 K provided both the largest BET and electrochemical surface areas, resulting in the highest photocurrent density of 2.2 mA cm^{-2} at 1.23 V_{RHE} for sunlight-driven water splitting. The photocurrent decreased upon increasing the oxynitride particle size. Although the smallest non-porous BaNbO_2N particles prepared *via* nitridation of the oxide calcined at 1073 K were highly crystalline, this did not contribute to increasing the surface reaction sites. This result demonstrates that the size reduction of oxynitride particles accompanied by an increase in the surface reaction area clearly enhances the water-splitting photoactivity. In fact, the photon flux in the solar spectrum (used to determine the number of charges generated during photoreactions) is constant depending on the light wavelength. Therefore, the activity result of perovskite BaNbO_2N depending on its particle size implies that the photo-response of the oxynitride is still relatively poor, and there is a large room for improvement in the activity up to the theoretical maximum STH efficiency of each oxynitride.

Besides the particle size, the surface shape of a perovskite or layered perovskite oxynitride also influences its photoactivity. As discussed previously, fluxes during nitridation can be used to readily change the shape of the oxynitride particles. The employment of different fluxes during nitridation and/or calcination of the starting oxide has been applied to vary the shape and size of 3D SrNbO_2N particles.⁹³ Moreover, the nitridation of $\text{La}_2\text{Ti}_2\text{O}_7$ using an NaCl-KCl flux produced highly porous brick-like LaTiO_2N particles.¹⁶ The surface shape of the oxynitride with a large surface area improved its photocatalytic water oxidation activity because of the high dispersion of CoO_x or IrO_2 cocatalysts over the large surface of the oxynitride. Meanwhile, rarely reported vertical SrNbO_2N nanorod arrays (as discussed earlier) with a 1D structure of the semiconductor promoted the strong separation of photogenerated charges exclusively in the vertical direction, thereby enhancing PEC activity.

Over the past few years, newly emerged 2D configurations of perovskite and layered perovskite oxynitrides have been found to provide a short diffusion transfer route for the photoexcited charges and thus lower their recombination.^{31,130–133} The electrons and holes photogenerated in $\text{HCa}_2\text{Nb}_3\text{O}_{10}$ nanosheets and selectively transferred to the surface of 2D $\text{Ca}_2\text{Nb}_3\text{O}_{10}^-$ plates without their self-recombination significantly improved photocatalytic water reduction activity under UV irradiation ($\lambda > 300$ nm).¹³⁴ A 2D layered perovskite $\text{K}_2\text{LaTa}_2\text{O}_6\text{N}$ was also stably active during the same photoreaction under visible-light illumination ($\lambda > 400$ nm).⁸⁹ The optical and electrical

properties and charge transport of 2D RP-type $\text{A}_2\text{BO}_3\text{N}$ ($\text{A} = \text{Ca}, \text{Sr}, \text{or Ba}$; $\text{B} = \text{Ta or Nb}$) layered perovskites have been studied based on density functional theory.¹³⁵ Thereby, the mobility of the electrons (or holes) photogenerated in the 2D layered perovskites was found to be much quicker than in the 3D perovskites. These results indicate that the advantages of 2D morphology (*i.e.*, the fast vertical charge transfer to the surface of the oxynitride plates) enhance water-splitting photoactivity.

Because the crystal structure of perovskite $\text{AB}(\text{O},\text{N})_3$ is cubic, the 3D shape of the oxynitride is typically synthesized. However, 2D-type starting oxides with a layered perovskite structure could be transformed into 2D oxynitrides capable of absorbing visible light. Fig. 14(B) shows the surface morphologies of the starting oxides and their corresponding oxynitrides, along with the visible-light-driven photoactivity of the 2D oxynitrides compared with the corresponding 3D type. In the upper row, it can be seen that the smooth layered LaKNaNbO_5 plates were converted to porous 2D LaNbON_2 *via* nitridation accompanied by volatilization of the K and Na species.¹²³ This conversion also worked well in the synthesis of counterpart LaTaON_2 using LaKNaTaO_5 .¹³¹ Interestingly, volatilization of the layered oxide along the [001] direction resulted in the shape of the oxynitride with exposed (010) surface facets remaining unchanged. The photocatalytic water oxidation activity of 2D LaNbON_2 with (010) facets was much higher than that of conventional 3D LaNbON_2 with mostly (100) facets. The efficient transfer of charges along the (010) facets of the 2D oxynitride and its low defect density resulting from the fast nitridation and the volatilization of the K and Na species led to its enhanced photoactivity. Meanwhile, layered perovskite Zr-doped $\text{Sr}_5\text{Nb}_4\text{O}_{15}$ with a 2D-type truncated octahedral structure was completely transformed to porous 2D-type SrNbO_2N particles with an identical morphology, as shown in the bottom row of Fig. 14(B).¹⁰⁵ The conversion process yielded oxynitrides with fewer defects, as discussed in detail in the earlier synthesis section using A-site cation-rich starting oxides. The 2D shape of the SrNbO_2N particles, as well as their high crystallinity, significantly heightened the PEC water-splitting activity of the oxynitride. The efficiency of charge separation by the 2D SrNbO_2N was much higher than that by the conventional 3D oxynitride. Moreover, the charge transfer resistance in the 2D oxynitride was relatively low. These results enable us to draw the same conclusion that oxynitrides with a 2D surface structure facilitate strong charge separation and fast transfer of charges photogenerated during water splitting.

4.3. Controlling the crystallographic facets

Crystal facet engineering for more efficient photoreaction activity by effectively promoting the preferential separation of charges between two adjacent anisotropic facets in a semiconductor during the reaction has been recently reported.^{27,136} Representatively, it has been demonstrated that monoclinic BiVO_4 particles with two different facets prepared *via* a hydrothermal method and the selective photo-deposition of cocatalysts could separate photogenerated holes in the (110) facets while collecting the counterpart electrons in the (010) facets.¹³⁷ Au, Pt, or Ag

nanoparticles were deposited on the (010) facets of BiVO_4 under light irradiation while MnO_x or PbO_2 particles were coated on the (110) facets of the oxide in the same manner. Both selective depositions of the cocatalysts, particularly the Pt and MnO_x particles, dramatically enhanced the photocatalytic water oxidation activity of the BiVO_4 particles. Similarly, spatial separation of the photogenerated holes and electrons with CoO_x and RhO_x cocatalysts, respectively, also improved the PEC water oxidation activity.¹³⁸ Moreover, optoelectrical simulation proved that facet engineering increased the efficiencies of not only charge

injection but also charge separation during the water oxidation reaction.

The facet engineering of cubic perovskite $\text{SrTiO}_3\text{:Al}$ was very effective at spatially separating the photogenerated charges to two different facets, thus driving overall water splitting *via* one-step photoexcitation.¹² $\text{Rh/Cr}_2\text{O}_3$ as an HER cocatalyst and CoOOH as an OER catalyst were sequentially photo-deposited onto the electron-attracting (100) facets and the hole-collecting (110) facets, respectively. The resulting anisotropic charge transport achieved high overall water-splitting activity that

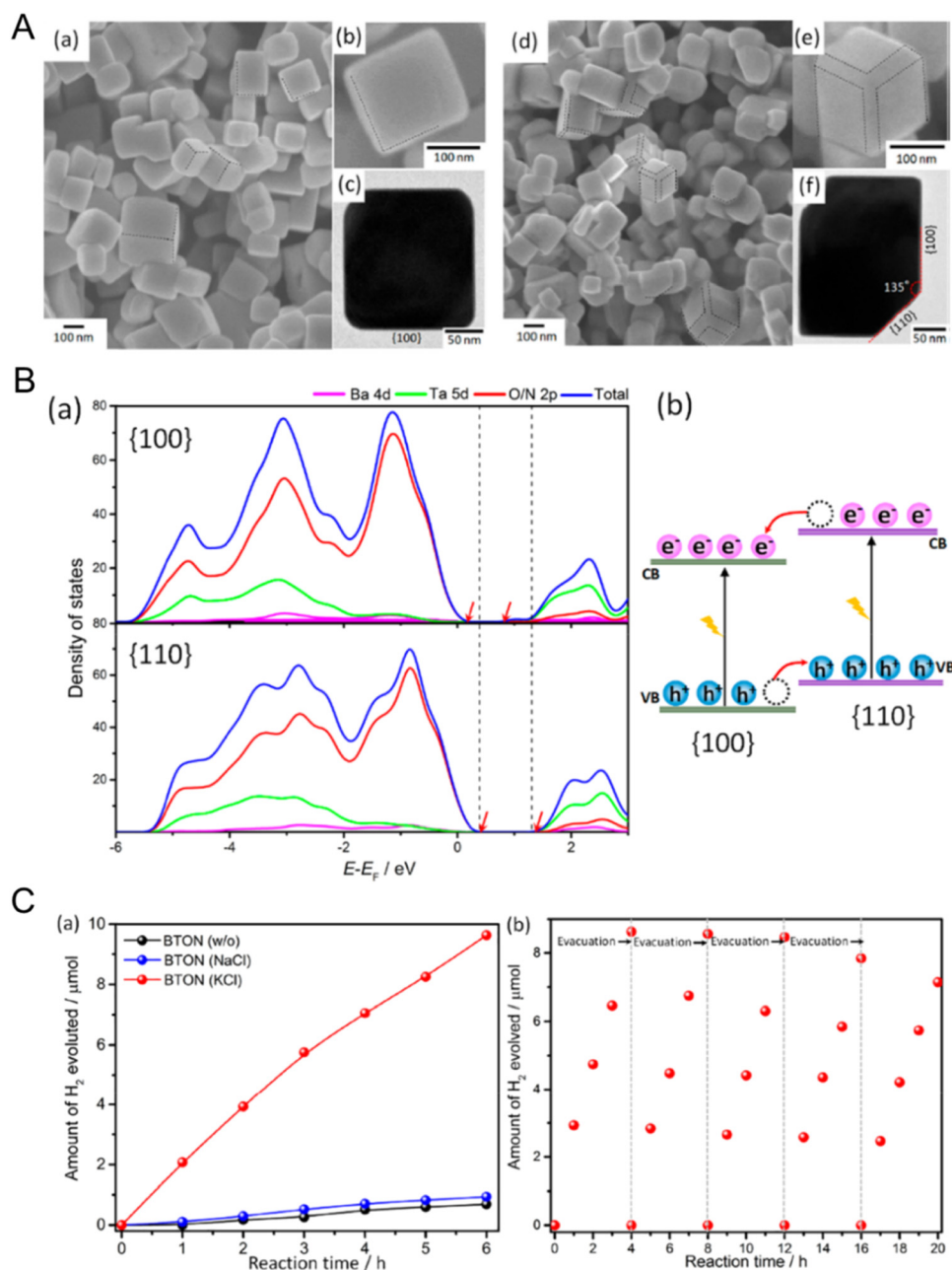


Fig. 15 Crystal facet engineering of perovskite and layered perovskite oxynitrides. (A) SEM and TEM images of BaTaO_2N particles prepared *via* flux-assisted nitridation using (a)–(c) NaCl and (d)–(f) KCl molten salts. (B) The density of states of the different energy levels of the (100) and (110) facets of BaTaO_2N prepared *via* KCl-assisted nitridation. (C) (a) Photocatalytic H_2 evolution activity using the two different BaTaO_2N particles and (b) cyclic testing using the oxynitride particles prepared *via* KCl-assisted nitridation. Reproduced with permission.¹³⁹ Copyright 2019, ACS.

was almost consistent with the theoretical quantum efficiency of SrTiO_3 . Unlike the oxide, it is difficult to prepare perovskite oxynitrides with exposed crystal facets to spatially separate charges because their crystal growth is not easily controlled during typical high-temperature nitridation. Fortunately, flux-assisted nitridation may offer a feasible means of tuning the surface morphology of oxynitrides, as discussed in the synthesis section. Symmetric BaTaO_2N particles with only the (100) and (110) facets exposed have been successfully prepared *via* flux-assisted nitridation, as displayed in Fig. 15(A).¹³⁹ The starting precursors of Ba and Ta were rearranged and then recrystallized in the molten salt during nitridation, which resulted in well-defined cubic BaTaO_2N particles of approximately 200 nm in size. Interestingly, the BaTaO_2N crystals prepared using an NaCl flux exhibited a cubic structure with six isotropic (100) facets whereas the oxynitride synthesized using a KCl flux presented a cubic-like structure with additional smooth (110) facets at the twelve corners of the (100) facets. The ionic species of fluxes become adsorbed on the crystal facets, thereby enabling faster growth and changes in the electrostatic interactions between the crystals. Because the ionic radius of six-coordinate K^+ (152 pm) is very similar to that of six-coordinate Ba^{2+} (149 pm), K^+ can more easily occupy the Ba^{2+} positions at the interface between BaTaO_2N and KCl. This not only suppresses the crystal growth of BaTaO_2N with the most stable (100) facets but also leads to the formation of secondary stable (110) facets. According to density functional theory calculations (Fig. 15(B)), the estimated surface energy of the (110) facets is higher than that of the (100) facets, meaning that the CBM and VBM potentials of the (110) facets are located

above those of the (100) facets. Thus, a facet junction is generated between the anisotropic (100) and (110) facets of a BaTaO_2N particle, thereby boosting the selective transfer of photogenerated electrons and holes to the (100) and (110) facets, respectively. Like the previously described BiVO_4 system, Pt and MnO_x particles were separately photo-deposited onto the (100) and (110) facets of BaTaO_2N . As a result, the BaTaO_2N crystals prepared using a KCl flux presented significantly improved water reduction activity compared with those using NaCl. Therefore, crystal facet engineering promoting selective separation and transfer of charges during photoreactions offers an effective strategy for enhancing the photoactivity of perovskite and layered perovskite oxynitrides by adjusting the crystal facets during their synthesis.

5. Conclusions and challenges

We discussed the preparation and use of perovskite $\text{AB}(\text{O},\text{N})_3$ and layered perovskite oxynitrides $\text{A}_{n+1}\text{B}_n(\text{O},\text{N})_{3n+1}$ as potential semiconductors for visible-light-driven artificial photosynthesis, as shown in Fig. 16. The materials are capable of absorbing visible light wavelengths up to 750 nm at most, and their band structures straddle the various redox potentials of the HER, OER, CER, H_2O_2 production, and CO_2 reduction, among others. Moreover, they are chemically stable in aqueous media over a wide pH range where photoreactions can take place. They fulfill the thermodynamic requirements for various photocatalytic processes to generate renewable, eco-friendly, and/or high value-added products such as H_2 and NH_3 with high theoretical

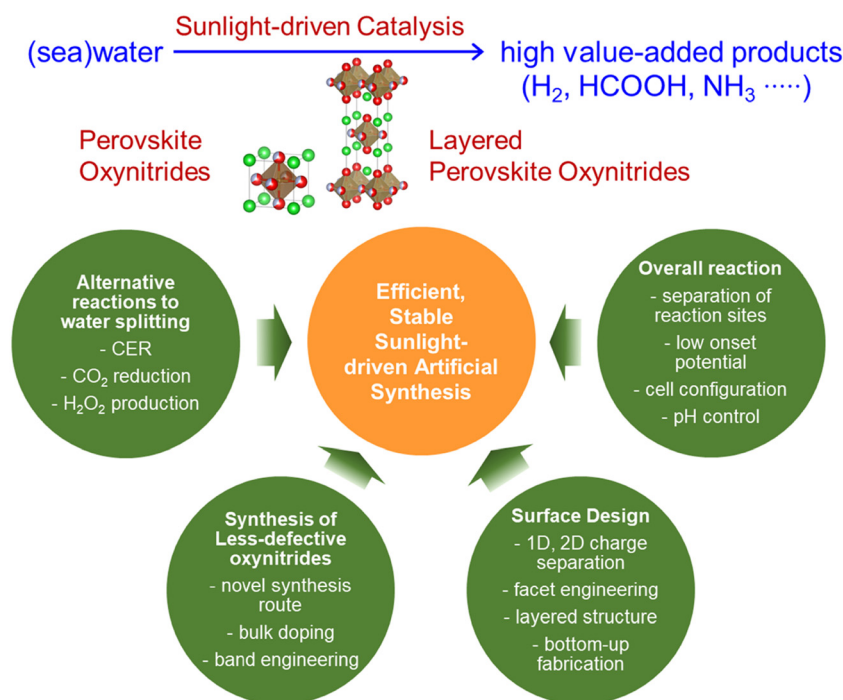


Fig. 16 Sunlight-driven catalysis and the challenges of perovskite and layered perovskite oxynitrides to artificially synthesize renewable, high value-added products such as H_2 .

solar-to-fuel energy conversion efficiency. Based on their favorable optical and chemical properties, perovskite and layered perovskite oxynitrides have been employed as both light absorbers and photocatalysts for visible-light-driven water splitting to produce H_2 over several decades. In addition, using them for CO_2 reduction and the CER as an alternative oxidation reaction to the OER to obtain high value-added products such as $HCOOH$ have recently been reported. The adaption of the CER, feasible by using seawater instead of fresh water, not only boosted the oxidation kinetics but also enhanced the H_2 production activity in neutral electrolytes. However, the perovskite and layered perovskite oxynitrides have shown low photoactivity and long-term stability during various photoreactions, which are still far from the requirements for commercialization (corresponding to STH values of above 10%).

Because Ti^{4+} , Ta^{5+} , and Nb^{5+} in the perovskite and layered perovskite oxynitrides are readily reduced during high-temperature nitridation, the surface and bulk properties of the latter can change due to the presence of defects and/or traces of impurities. Defects in perovskite and layered perovskite oxynitrides boost the recombination of photogenerated holes and electrons during photoreactions, thereby limiting their photoactivity. This phenomenon can be reduced by various synthesis strategies and surface and bulk engineering including doping and substitution. Moreover, photooxidation of the oxynitride surface, which results in the release of N_2 and a positive shift in the surface Fermi level, can be completely suppressed *via* surface modification to provide separated compartments for HER and OER cocatalysts or passivation layers.

Controlling the surface morphology of the oxynitride particles and/or electrodes can spatially separate the transfer orientation of photogenerated holes and electrons, thus leading to dramatic improvement in the sunlight-driven catalytic activity. Although 1D and 2D perovskite $AB(O,N)_3$ were rarely studied in terms of the synthesis method, the morphologically modified surfaces of the oxynitrides were found to largely increase the separation efficiency of charges generated during PEC water splitting. It is known that the surface shape of metal oxynitrides such as Ta_3N_5 is relatively easy to be transformed even through high-temperature nitridation. However, it is difficult to prepare $AB(O,N)_3$ with 1D or 2D surface morphology because two different cations at A- and B-sites are typically sintered at the high temperature probably causing the change in the surface structure. Alternatively, the charges generated during photoreactions can be spatially separated by crystal facet engineering and the employment of layered perovskite oxynitrides. The crystal facet engineering of $BaTaO_2N$, materialized by flux-assisted nitridation including a recrystallization stage, proved that (100) and (110) facets play roles as electron-attracting and hole-collecting sites, respectively. The layered perovskite oxynitrides with a 2D crystal structure, which allow selective transfer of holes and electrons in crystal lattices intrinsically, can be also utilized for sunlight-driven catalysis. The spatial charge separation in photoelectrodes, as well as on oxynitride particles, is likely to be necessary for efficient, stable photoreactions. In particulate photoelectrodes, the electron transfer from

oxynitride particles to a conductive substrate is significantly resistive due to the loose adhesion between them. The bottom-up fabrication to directly grow oxynitride crystals on the conductive substrate can be a solution to obtain a fast electron pathway comparable to the hole transfer rate to surfaces of oxynitrides. Therefore, both the novel synthesis of less-defective oxynitrides to reduce the recombination of photogenerated charges and the surface design of the oxynitrides to make spatially selective separation and transfer of the charges must be challenges to achieving efficient, stable sunlight-driven artificial synthesis using perovskite and layered perovskite oxynitrides.

The overall water splitting of perovskite and layered perovskite oxynitrides by one-step photoexcitation can be the most ideal way to facilitate the low-cost production of H_2 gas on a large scale. The overall water splitting of a $ATaO_2N$ ($A = Sr, Ba$) particle has been demonstrated; however its activity is still limited. The separation of both HER and OER sites at the oxynitride surface is likely to be necessary to drive water splitting, while HER sites with a cocatalyst were distinguished from the surface of oxides such as $SrTiO_3$. The selective deposition of the cocatalyst and crystal facet engineering of the oxynitrides are challenging for efficient overall water splitting. In a PEC system, the anodic photoresponse of perovskite and layered perovskite oxynitrides should be below 0 V_{RHE} based on their flat band potentials (which are negative). However, the actual photocurrent onset is limited to the range of 0.5 to 0.6 V_{RHE} . Unassisted solar water splitting *via* PEC cells comprising perovskite and layered perovskite oxynitrides combined with other cathode materials has rarely been reported. Thus, efficient photoanodes must be able to generate significant photocurrent density at a moderate potential (0.4 to 0.6 V_{RHE}) based on the overlap with the photocathode during overall water splitting. It is necessary to apply various approaches to improve their water-splitting activity by, for instance, optimizing their optical and physical properties. Moreover, a better understanding of the effect of pH (photoanodes are more active at higher pH values while photocathodes work well at lower pH values) must also be achieved to obtain a high rate of H_2 production. Therefore, these challenges are directly related to the photoactivity and long-term stability of these materials for efficient artificial photosynthesis.

Conflicts of interest

The authors have no conflicts of interest relevant to this study to disclose.

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