Nano-Micro Letters



Cite as Nano-Micro Lett. (2022) 14:78

Received: 17 December 2021 Accepted: 9 February 2022 Published online: 25 March 2022 © The Author(s) 2022

HIGHLIGHTS

Multilayer Strategy for Photoelectrochemical Hydrogen Generation: New Electrode Architecture that Alleviates Multiple Bottlenecks

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- A multilayer architecture of layers with different functions alleviates bottlenecks in photoelectrochemical (PEC) hydrogen generation. Precise thickness control within a few nanometers defines each layer's functionality.
- A Bi₂S₃/NiS/NiFeO/TiO₂ photoanode had a photocurrent density of 33.3 mA cm⁻² at 1.23 VRHE under AM 1.5 G illumination.
- Noble-metal-free seawater splitting was performed in an integrated PEC-electrocatalytic cell with an NiS electrocathode and Bi₂S₃/ NiS/NiFeO/TiO₂ photoanode.

ABSTRACT Years of research have demonstrated that the use of multiple components is essential to the development of a commercial photoelectrode to address specific bottlenecks, such as low charge separation and injection efficiency, low carrier diffusion length and lifetime, and



poor durability. A facile strategy for the synthesis of multilayered photoanodes from atomic-layer-deposited ultrathin films has enabled a new type of electrode architecture with a total multilayer thickness of 15–17 nm. We illustrate the advantages of this electrode architecture by using nanolayers to address different bottlenecks, thus producing a multilayer photoelectrode with improved interface kinetics and shorter electron transport path, as determined by interface analyses. The photocurrent density was twice that of the bare structure and reached a maximum of 33.3 ± 2.1 mA cm⁻² at 1.23 V_{RHE}. An integrated overall water-splitting cell consisting of an electrocatalytic NiS cathode and Bi₂S₃/NiS/NiFeO/TiO₂ photoanode was used for precious-metal-free seawater splitting at a cell voltage of 1.23 V without degradation. The results and root analyses suggest that the distinctive advantages of the electrode architecture, which are superior to those of bulk bottom-up core–shell and hierarchical architectures, originate from the high density of active sites and nanometer-scale layer thickness, which enhance the suitability for interface-oriented energy conversion processes.

KEYWORDS Atomic layer deposition; Bismuth sulfide; n-p junction; Photoelectrochemical; Nickel sulfide

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1 Introduction

The generation of hydrogen, an ideal fuel, using photoelectrochemical (PEC) water splitting can establish a clean green energy source and alleviate the increasing energy crisis [1, 2]. The production of hydrogen by PEC water splitting is a subject of cutting-edge research that can ensure the stability of our ecosystem [3–5]. However, low solar-to-hydrogen (STH) efficiency and poor durability are the main barriers to the commercialization of the PEC electrodes reported to date [6, 7]. In addition, hydrogen generation should use nontoxic and Earth-abundant components to ensure affordability [8]. Highly active precious metal catalysts (such as IrO₂ and Pt) are not appropriate candidates for replacing nonrenewable energy sources; the use of toxic elements requires more costly processing and disposal to maintain safety [9]. A light-absorbing semiconductor and surface catalyst are the main components of a PEC electrode. However, research has shown that several additional components, such as a heterojunction for effective charge separation and passivation layers for longer life, are essential to the development of commercial photoelectrodes [10–13]. Therefore, an appropriate choice of each component is critical for the full utilization of solar energy and for obtaining an efficient yield of light-induced charge carriers. Choosing narrowband-gap semiconductors increases the likelihood of high charge carrier generation by extended light absorbance of solar spectrum. The formation of heterojunctions with opposite semiconductors is intensively studied due to the natural advantage of immense charge separation and long charge carrier lifetime. Furthermore, photocurrent generation is an interface-dominant electrochemical process that requires an extremely high density of surface-active sites, which accelerate the interface reactions to a sufficiently high rate. However, Effects of surface states (SS) are unavoidable, and surface recombination through SS greatly affects the net photocurrent at electrode-electrolyte interfaces [14-17]. Surface recombination is responsible for more than 90% of the loss of light-induced charge carriers in various photoelectrodes. The integration of surface co-catalysts is the conventional method of alleviating the detrimental effects of SS and trapping charges at the interface before recombination. Therefore, the design of a co-catalyst within the thickness of the depletion region helps in an accurate analysis of the adaptive junctions that coexist with SS and the co-catalyst. Finally, passivation layers are essential for meeting industrial durability standards in large-scale applications.

Motivated by the above considerations, we designed a multilayer photoanode model in which each layer plays a different role, specifically, the primary light absorber, n-p junction, cocatalyst, and passivation layer, to obtain the maximum photocurrent output. The fabrication of the multilayer structure involves engineering difficulties such as the need to preserve the lower layers, selecting compatible synthesis methods, the buildup of solid-solid junction resistance, and, most importantly, avoiding the masking of primary light absorber. Therefore, we adopted atomic layer deposition (ALD) as the main tool to produce the multilayer structure because it can produce conformal thin films over high-aspect-ratio structures [18–20]. The dry, inert and high vacuum operating conditions of ALD make it possible to eliminate the engineering difficulties mentioned above in order to form multilayer electrodes as thin as possible. This model electrode differs from classical core-shell and hierarchical structures because it ensures multilayer formation within the band-bending region (5-10 nm) and, consequently, yields the fastest charge transfer kinetics. The use of atomically thin multilayers can decrease the migration distance of photogenerated charge carriers and increase the probability of carrier transfer at electrode-electrolyte interfaces compared with that of the bulk counterpart [21].

Regarding material selection, in addition to well-studied inorganic metal oxides with band gaps above 2.0 eV, metal chalcogenides have recently shown great potential for hydrogen generation because of their large light absorption [22]. In particular, n-type bismuth sulfide (Bi_2S_3) , which has a direct narrow band gap (1.30-1.35 eV), can harvest the full UV-visible (UV-vis) region of the solar spectrum [23, 24]. NiS (0.9–1.0 eV) was then selected as its p-type counterpart because its band edge positions are compatible with those of several well-developed semiconductors, including bismuth sulfide, and it is thus suitable for forming heterojunctions [25, 26]. The cascade alignment of band edge positions between bismuth sulfide and nickel sulfide semiconductors can produce photogenerated carriers moving in opposite directions, which result in high photocurrent and STH efficiency. The inherent problem of the low bandgap semiconductor is photocorrosion and could be alleviated by adopting a co-catalyst to transfer the accumulated charge carriers to the electrolyte before it oxidizes the primary light absorber. Transition metal oxides/ hydroxides/oxyhydroxides exhibit the best catalytic activity in PEC water splitting [18, 27]. In particular, nickel- and ironbased oxyhydroxides are widely reported to exhibit successful charge tunneling at the interface. Our group recently reported the stoichiometric dependence of the NiFeOOH co-catalyst on hematite photoanodes in the charge tunneling process [18]. At the top of the electrode, a thin layer of titanium dioxide was used as a passivation layer to avoid direct contact of the electrolyte with the active components [28].

In this study, the $Bi_2S_3/NiS/NiFeO/TiO_2$ multilayer photoanode was systematically analyzed to identify the role of each functional layer deposited on bismuth sulfide. In addition, the mechanism of inherent SS passivation and ladderlike hole transport from the valence band (VB) of bismuth sulfide to the electrolyte was elucidated through interface studies. Furthermore, seawater splitting by a precious-metalfree integrated PEC-electrocatalytic (EC) water-splitting cell with a $Bi_2S_3/NiS/NiFeO/TiO_2$ photoanode and NiS electrocathode was demonstrated.

2 Experimental

2.1 Fabrication of Bi₂S₃/NiS/Ni_{0.75}Fe_{2.25}O₄/TiO₂ Photoanodes

2.1.1 Synthesis of Bi₂S₃ Photoanodes

Bismuth sulfide nanostructures were grown by a previously reported two-step solvothermal process [29]. First, 0.05 M tin chloride and 0.1 M thioacetamide are dissolved in 40 mL of ethanol and maintained at 80 °C for 4 h with a cleaned fluorine-doped tin oxide (FTO) glass substrate in an air-tight reactor. The formed SnS_x films over FTO substrates were cleaned several times with ethanol and dried under N₂ flow. Then, the SnS_x films were hydrothermally treated with a 0.04 M bismuth chloride acidic solution for 24 h at 100 °C to form Bi_2S_3 . The dried samples were annealed under N₂ atmosphere at 450 °C for 2 h.

2.1.2 Synthesis of Bi₂S₃/NiS Photoanodes

NiO thin films were deposited using a homemade flow-type ALD reactor maintained at 175 °C and 800 mTorr. Argon (99.999%) was used as both the carrier (50 sccm) and purging (250 sccm) gas. Nickel cyclopentadienyl [Ni(Cp)₂] (Sigma-Aldrich, USA) and ozone (5% in O_2) were used in the ALD

process as the nickel precursor and counter-reactant, respectively [30]. A single ALD cycle consisted of four steps: a 1.5 s Ni(Cp)₂ pulse, a 30 s precursor purge, a 3 s ozone pulse, and a 60 s ozone purge. Before ALD was performed on the Bi₂S₃ nanostructures, the ALD conditions were optimized using single-crystalline p-type silica (100) flat substrates (LG Siltron, Korea). A dry sulfurization process was employed to drive the anion exchange reaction (AER) to convert NiO into NiS [31]. To obtain Bi₂S₃/NiS photoanodes, NiO thin films with various thicknesses were coated on the Bi₂S₃/FTO substrates; thermal vulcanization was then conducted by placing 2 g of S powder and the FTO/Bi₂S₃/NiO samples in the upstream and downstream parts of a tube furnace, respectively, and heating them to 350 °C under N₂ atmosphere for 1 h.

2.1.3 Synthesis of Bi₂S₃/NiS/Ni_{0.75}Fe_{2.25}O₄ Photoanodes

Atomic-layered $Ni_{0.75}Fe_{2.25}O_4$ (NiFeO) was deposited on the Bi_2S_3 /NiS photoanodes following a procedure similar to that reported in our previous work [18]. The NiFeO thickness was controlled by varying the number of ALD supercycles.

2.1.4 Synthesis of Bi₂S₃/NiS/Ni_{0.75}Fe_{2.25}O₄/TiO₂ Photoanodes

Titanium oxide was coated on the prepared $Bi_2S_3/NiS/NiFeO$ samples via ALD using a homemade ALD system at 175 °C using tetrakis(dimethylamido)titanium (99.9%, UP Chemicals) and hydrogen peroxide (30 vol% in water, Sigma-Aldrich) as the reactants [32]. The thickness of the TiO₂ films was controlled by varying the number of ALD cycles. The coated films were used directly for PEC analysis without any postheat treatment processes.

2.1.5 Synthesis of FTO/NiS Electrodes

NiO thin films (10 nm) were deposited on cleaned bare FTO substrates. A dry sulfurization process was then applied to drive the AER to convert NiO into NiS.

2.2 Characterization

The crystallinity of each sample was analyzed using high-resolution X-ray diffraction (XRD) measurements

(PANalytical) with a 3D-PIXcel detector and Cu Ka radiation at 60 kV and 55 mA. High-resolution X-ray photoelectron spectroscopy (XPS) was employed using Ka radiation and seven Channeltron detectors. The structural and morphological properties of the electrodes were studied using high-resolution scanning electron microscopy (HR-SEM; JEOL JSM-7500F) coupled with energy-dispersive X-ray spectroscopy at an acceleration voltage of 15 kV, and high-resolution transmission electron microscopy (HR-TEM, TECNAI G2 F20). UV-vis spectroscopy (VARIAN) was also used to observe the optical properties of the photoanodes using bare FTO as a reference. A time-resolved photoluminescence (TRPL) study was performed using an inverted-type scanning confocal microscope (MicroTime-200, Picoquant, Germany) with a $40 \times$ objective. A single-mode pulsed diode laser (LDH-P-C-470, Picoquant, Germany) with a ~ 30 ps pulse width and 0.3 mW average power was used as an excitation source. A dichroic mirror (490 DCXR, AHF), two long-pass filters (HQ500lp, AHF; FEL600, Thorlabs), and a single-photon avalanche diode (PDM series, MPD) were used to collect the emission from the samples. Time-correlated single-photon counting was used to count the fluorescence photons. Fitting was done using a biexponential decay model.

2.3 Photoelectrochemical Measurements

The PEC measurements were performed in the presence of 0.25 M Na₂S (pH~12.5) at~25 °C. Ag/AgCl and Pt sheets were used as reference and counter electrodes, respectively. The measured potentials versus Ag/AgCl were converted to reversible hydrogen electrode (RHE) using the Nernst equation. A 300 W Xe lamp was used as the light source for an AM 1.5 G simulation, and the incident power density (100 mW cm⁻²) was calibrated using a standard silicon cell. The open-circuit voltage decay was measured using the same setup. Photoelectrochemical impedance spectroscopy (PEIS) measurements were made at an amplitude of 10 mV and frequencies ranging from 10^{-2} to 10^{6} Hz. Mott–Schottky analysis was conducted under dark conditions. The gas evolution during photolysis was analyzed using gas chromatography (074-594-P1E Micro GC Fusion, INFICON) in 0.25 M Na₂SO₄ (pH ~ 7.0) electrolyte.

3 Results and Discussion

3.1 Physical Characterization of Bi₂S₃/NiS/NiFeO

The XRD diffractograms of as-prepared Bi₂S₃ and Bi₂S₃/ NiS are shown in Fig. 1a. All the diffraction peaks matched those of orthorhombic bismuthinite (JCPDS No. 017-0320), and no impurity phases were identified [23]. Furthermore, the sharp peaks that were observed in the diffraction pattern after the formation of NiS is due to annealing in the S environment indicated that the prepared Bi₂S₃ was very crystalline. No diffraction peaks corresponding to the NiS phase were observed due to the predominance of the large Bi_2S_3 core (~2 µm) over the NiS nanolayer (~5 nm). The surface composition of the Bi₂S₃/NiS/NiFeO photoanode was analyzed using XPS; peaks for the spin orbitals Bi $4f_{7/2}$ and Bi 4f_{5/2} appear at 158.7 and 164.0 eV, respectively, with a splitting energy of 5.3 eV (Fig. 1b) [33]. Two additional peaks corresponding to the S $2p_{3/2}$ and S $2p_{1/2}$ spin orbitals at 161.36 and 162.9 eV, respectively, overlap the Bi 4f peaks [34]. Figure 1c shows the high-resolution Ni 2p spectra, which indicate that most of the Ni atoms are in the +2 oxidation state, as shown by the intense peak at 856.29 eV and its satellite peak at 861.50 eV. The peak at 859.13 eV indicates the coexistence of Ni³⁺ ions in the NiFeO co-catalyst layer. Figure 1d shows the high-resolution Fe 2p spectra, where the +2 and +3 oxidation states are represented by the intense peaks at 710.19 and 712.16 eV, respectively. The UV-vis absorption spectra of Bi₂S₃ and Bi_2S_3/NiS are shown in Fig. 1e, where both photoanodes exhibit strong light absorption in the full visible wavelength range because of the intrinsic narrow band gap of Bi₂S₃. There is a slight increase in light absorption, which was not significantly altered by increasing the thickness of the NiS nanoshell because the nanoshell thickness is negligible compared to the Bi₂S₃ core width. To further confirm the presence of the NiS phase, a detailed physical characterization of FTO/NiS is conducted and presented in Fig. S1. The absorbance is compared in Fig. S2a. A sharp increase in absorption appeared at 260 nm, and the absorption range covered the entire visible light region [35]. The Tauc plot of NiS indicates a band gap of approximately 1.01 eV, which is an intrinsic characteristic of NiS, as shown in Fig. S2b. The Tauc plots of Bi2S3 and Bi2S3/NiS for direct allowed transitions revealed a band gap of ~ 1.32 eV, as shown in Fig. 1f.



Fig. 1 a XRD spectrum of Bi_2S_3 and Bi_2S_3 /NiS photoanodes. High-resolution XPS spectra of Bi_2S_3 /NiS/NiFeO photoanode: **b** Bi 4f and S 2p, **c** Ni 2p, and **d** Fe 2p. **e** UV–vis spectra of Bi_2S_3 and Bi_2S_3 /NiS photoanodes measured with bare cleaned FTO as a reference to eliminate absorption by air and FTO. **f** Tauc plots of Bi_2S_3 and Bi_2S_3 /NiS for direct allowed transitions

Figure 2a-c shows HR-SEM images of bismuth sulfide before and after the addition of the NiS layer (by the ALD and subsequent sulfurization of a 5 nm NiO layer). As shown in Figs. 2a and S3, the bismuth sulfide rods are vertically aligned and have a height and width of 10 and 1 μ m, respectively. Magnified images of the bare and NiS-coated bismuth sulfide rods are shown in Fig. 2b-c, respectively; the drastic increase in surface roughness originates from the NiS nanoshell. The interface properties and the structural changes resulting from anion exchange were examined by HR-TEM analysis. An HR-TEM image of the bare bismuth sulfide rod is shown in Fig. 2d, and the ordered bright spots in the selected area electron diffraction (SAED) pattern indicate the high crystallinity and orientation of the bismuth sulfide rods, as shown in Fig. 2e. The higher-magnification image in Fig. 2f shows lattice fringes of 0.36 nm, which correspond to the (130) crystal plane of orthorhombic bismuth sulfide [24, 36]. The bismuth sulfide rod coated with the 5 nm ALDdeposited NiO film is shown in Fig. 2g. The uniformity of the NiO film demonstrates that ALD can be used to coat high-aspect-ratio nanostructures with excellent uniformity and conformality [37]. After the AER, a NiS shell with a thickness of 6-7 nm can be observed on the bismuth sulfide

rods, as shown in Fig. 2h. The fast Fourier transform (FFT) image with ordered bright spots (inset in Fig. 2i) also confirms the crystalline nature of the shell material; thus, the NiS nanoshell can be expected to have an appropriate band gap and alignment with bismuth sulfide. The higher-magnification images in Fig. 2j–k reveal highly ordered lattice fringes with a *d* spacing of 0.22 nm, which corresponds to the (211) crystal plane of the NiS phase (1:1 stoichiometry) [38]. The elemental distributions of the Bi₂S₃/NiS/NiFeO photoanode (Fig. 2l–o) confirm the discrete distributions of elemental Ni and Fe over the bismuth sulfide rods, which clearly indicates continuous shell formation. Features corresponding to NiFeO are not visible because the deposited NiFeO films are ultrathin. The optimized NiFeO thickness, as indicated by the electrochemical performance, is 0.6–1.0 nm.

3.2 Photoelectrochemical Performance

The $Bi_2S_3/NiS/NiFeO/TiO_2$ photoanode has three functional layers with a total thickness of 15–17 nm, which directly affect the electrochemical interfacial kinetics. The NiS nanoshell is the first functional nanoshell/nanolayer over the Bi_2S_3 primary light absorber. Both bismuth sulfide and



Fig. 2 HR-SEM images of **a**, **b** Bi_2S_3 and **c** Bi_2S_3/NiS . **d** HR-TEM image of Bi_2S_3 and **e** SAED pattern of the area in the green square in **d**. **f** Diffraction plane analysis of Bi_2S_3 . HR-TEM images of **g** Bi_2S_3/NiO and **h-k** Bi_2S_3/NiS (inset: FFT image). **l-o** Elemental mapping of $Bi_2S_3/NiS/NiFeO$ photoanode

nickel sulfide are narrow-band-gap semiconductors with opposite majority charge carriers, and they tend to form type-II cascade heterojunctions in equilibrium [39, 40]. The p-type behavior of NiS is confirmed by the *J*–*T* curve of the FTO/NiS photocathode at 0.0 V_{RHE} under chopped illumination (Fig. S4) [35]. Scheme 1 shows the band edge positions and the electrochemical potentials of possible interface reactions with respect to RHE at pH~7. The

 Bi_2S_3/NiS photoanode generated a photocurrent density of 27.4 ± 1.5 mA cm⁻² at 1.23 V_{RHE}, which is more than twice that of the bare Bi_2S_3 photoanode, as shown in Fig. 3a. The nonzero dark current represents the formation of sulfite, as shown in Sch. 1. The sharp strike current of Bi_2S_3 and Bi_2S_3/NiS indicates a lack of surface activity and loss of charge carriers through surface recombination. When the second functional layer, the NiFeO co-catalyst, was added,



Scheme 1 Band diagram of heterojunction, with band edges and chemical redox potentials of possible reactions with respect to RHE

the strike current disappeared, and the photocurrent density increased to 33.3 ± 2.1 mA cm⁻² as a result of accelerated charge transfer from the VB of NiS to the electrolyte. The PEC performance of the Bi₂S₃/NiS/NiFeO photoanode is compared with the performance reported in recent studies in Table S1. Each nanoshell thickness is crucial to PEC water splitting because the individual electrocatalytic activity of NiS should not fade out the PEC activity of Bi₂S₃. The effect of nanoshell thickness was evaluated using *J*–*V* curves obtained under chopped illumination. As shown in Fig. S5, the NiS shell prepared from a 5-nm-thick NiO shell generates the maximum photocurrent density. The dark current increases with NiS shell thickness, indicating that

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Fig. 3 a J-V curve measured under AM 1.5 G illumination, **b** open-circuit decay curve, **c** ABPE analysis, **d** IPCE analysis, **e** Mott–Schottky plot using double-layer capacitance measured by dark EIS, **f** charge separation efficiency, and **g** charge injection efficiency. **h** Nyquist plot of photoanodes at 1.23 V_{RHE}

the NiS electrocatalyst is becoming independent, which reduces the photocurrent density. The total current density (dark + light) across the NiS/electrolyte interface was compared with the charge transfer resistance (R_{ct}) as shown in Fig. S5e. The R_{ct} increases slightly with increased shell thickness (10 nm) [41]. Therefore, the shell thickness was optimized as 5 nm, based on minimum R_{ct} and maximum photocurrent [31]. Furthermore, a control sample was synthesized by depositing a bulk NiS layer on Bi₂S₃ using the SILAR (Successive Ion Layer Reaction and Adsorption) method. The bulk NiS layer completely masks the PEC behavior as shown in Fig. S6. The effect of NiFeO thickness was evaluated using atomic thicknesses of 0.6, 1, 2, and 3 nm. The results (Fig. S7) indicate that a thickness of 1 nm is optimal for both catalytic activity and interfacial kinetics. The R_{ct} with respect to the different thicknesses of NiFeO follows a trend similar to that of the thickness of the NiS layer [42]. The open-circuit potential (OCP) decay curves in Fig. 3b show that $V_{\rm ph}$ significantly improved with the addition of the NiS and NiFeO functional layers. After the light source is turned off, the photoinduced electrons recombine with the holes or become trapped by SS [43]. The decay rate of the OCP to its dark equilibrium condition represents the life time of photoinduced electrons [44]. $Bi_2S_3/$ NiS/NiFeO exhibits the slowest decay among the photoanodes, indicating reduced non-radiative recombination. The negative-shifted OCP of Bi₂S₃/NiS under dark condition is due to the detrimental potential drop across the Helmholtz layer [45]. The NiFeO co-catalyst incorporation successfully alleviated the potential drop and maintained the OCP at its original value. The photocurrent onset (V_{on}) is a key indicator of surface catalyst efficiency; here, the 170 mV cathodic shift of Bi₂S₃/NiS after the addition of the NiS nanoshell originated from the intrinsic catalytic activity of Ni-S active sites (Fig. S8a) [11]. The NiFeO co-catalyst provided a shift of ~250 mV in addition to that of Bi₂S₃/NiS; thus, the V_{on} value of $Bi_2S_3/NiS/NiFeO$ is $-0.25 V_{RHE}$. V_{on} is generally influenced by two factors, the photovoltage (V_{ph}) and kinetic overpotential (η), as follows: $E_{\text{redox(dark)}} - V_{\text{on}} = V_{\text{ph}} - \eta$, where $E_{\text{redox}(\text{dark})}$ is the electrochemical potential of the electrolyte under dark [46]. The obtained overall cathodic shift (~420 mV relative to V_{on} of bare Bi₂S₃) is lower than the increase in $V_{\rm ph}$ (~70 mV) measured under equilibrium conditions, as shown in Fig. 3b. Therefore, the obtained cathode shift can mainly be attributed to the decrease in kinetic overpotential as shown in Fig. S8b and Table S3. As shown

in Fig. 3c, the highest applied bias photon-to-current efficiency (ABPE) is observed for Bi2S3/NiS/NiFeO (12.99% at 0.64 V_{RHE}), followed by Bi₂S₃/NiS (6.56% at 0.80 V_{RHE}) and Bi_2S_3 (4.48% at 0.82 V_{RHE}). Figure 3d shows the incident photon to current conversion efficiency (IPCE) of bismuth sulfide photoanodes. The order of electrodes in terms of IPCE is as follows: Bi₂S₃/NiS/NiFeO (76.78%) > Bi₂S₃/NiS $(52.16\%) > Bi_2S_3$ (36.16%) at 450 nm, and this corresponds to the order seen in the J-V curves. The similar shape of all IPCE spectra in the ultra-violet and visible region indicates the identical light absorption in corresponding region. The low IPCE in the ultra-violet and blue region represents the surface recombination effect. The declining IPCE response from the green to red region denotes the low diffusion length of charge carriers. The elevated IPCE response at the infrared region originates mainly from the photo absorbance of NiS in Bi₂S₃/NiS and Bi₂S₃/NiS/NiFeO photoanodes.

Mott-Schottky analysis was conducted in dark conditions, and the carrier density was found to be 2.87×10^{21} and 20.58×10^{21} cm⁻³ for Bi₂S₃ and Bi₂S₃/NiS, respectively, as shown in Fig. 3e. This drastic enhancement provides strong evidence for increased charge separation at the interface due to n-p junction formation. The addition of the NiFeO co-catalyst also increases the carrier concentration by~3.5 times because the rapid tunneling of photogenerated charges through NiFeO reduces the probability of charge recombination. The yield of photogenerated charges at the interface (η_{bulk}) for different Bi₂S₃ and Bi₂S₃/NiS samples was calculated using the procedures described in previous reports (Fig. S9). The calculated values of η_{bulk} for all the samples are plotted in Fig. 3f. The η_{bulk} value of Bi₂S₃/NiS is greater than that of Bi₂S₃ across the potential window. In particular, the more significant difference in η_{bulk} at higher potentials (>0.9 V) indicates that NiS contributes mainly to charge separation, not tunneling. Although NiS is a wellknown oxygen evolution reaction catalyst, the low η_{bulk} at lower potentials and large strike currents indicate the presence of SS on NiS, which is also reflected in the low $\eta_{surface}$, as shown in Fig. 3g. The NiFeO co-catalyst increases η_{bulk} , which reaches 100% at 1.2 V_{RHE} by reducing charge recombination through rapid charge removal from the VB of NiS. The larger increase in η_{surface} at lower potentials (<0.6 V_{RHE}) when the NiFeO co-catalyst is present demonstrates accelerated charge tunneling at the NiFeO-electrolyte interface. Nyquist plots of the Bi₂S₃, Bi₂S₃/NiS, and Bi₂S₃/NiS/NiFeO photoanodes under illumination were obtained at 1.23 V_{RHE},

as shown in Fig. 3h. The drastic decrease in R_{ct} indicates accelerated charge tunneling at the NiS–electrolyte and NiFeO–electrolyte interfaces for the Bi₂S₃/NiS and Bi₂S₃/NiS/NiFeO photoanodes, respectively.

3.3 Charge Transfer and Recombination Kinetics

The charge carrier kinetics were studied through the PEIS measurements with linear bias interval. The Nyquist plot of $Bi_2S_3/NiS/NiFeO$ at 0.30 V_{RHE} has only two semicircles, which correspond to the capacitance of the bulk space-charge region/Helmholtz layer (C_{bulk}) and charge tunneling ($C_{SS/cat}$). Both the Bi_2S_3 and Bi_2S_3/NiS photoanodes show

three semicircles, including C_{trap} in the medium-frequency range, which can be attributed to reaction intermediates or trap surface states (Fig. 4a). The disappearance of C_{trap} in Bi₂S₃/NiS/NiFeO indicates the successful passivation of surface traps by the NiFeO co-catalyst [47]. The R_{ct} values were calculated from the Nyquist plot by fitting the data using the equivalent circuit (in Fig. 4d), as shown in Fig. 4b. Bi₂S₃ shows the highest R_{ct} throughout the potential window. This result highlights the high density of surface traps on the Bi₂S₃ surface, which causes high charge recombination and thus low $\eta_{surface}$; the photogenerated charges can overcome the trapping barrier only at high applied potentials. The addition of NiFeO decreases R_{ct} by two orders



Fig. 4 a Nyquist plot of bismuth sulfide photoanodes at 0.30 V_{RHE} , **b** interface charge transfer resistance (R_{cl}), and **c** distribution of $C_{SS/cat}$ and N_{SS} calculated from PEIS measurements at various potentials under illumination in 0.25 M Na₂S electrolyte, **d** Electronic circuit used to fit the PEIS data, **e** charge recombination rate constant (k_{rec}), **f** TRPL transients, where solid blue line represents a bi-exponential fitting (table: fitted values), **g** charge transfer rate constant (k_{cl}), and **h** overall quantum efficiency of photoanodes

of magnitude, demonstrating accelerated charge tunneling through NiFeO. $C_{SS/cat}$ decreases gradually with increasing bias for all the photoanodes owing to successful band bending and the release of accumulated charges to the electrolyte (Fig. 4c). The SS distribution was calculated from $C_{SS/cat}$ as a function of applied potential (*E*) as follows [48]:

$$N_{\rm ss}(E) = \frac{C_{\rm SS/cat}(E)}{q} \tag{1}$$

where q is the electron charge $(1.602 \times 10^{-19} \text{ C})$.

The N_{SS} distribution is dominant for the Bi₂S₃/NiS/NiFeO photoanode because of the increased carrier density at the NiFeO/electrolyte interface, which is identical to the carrier density measured by Mott-Schottky analysis. The shift in the maximum $N_{\rm SS}$ coincides with $V_{\rm on}$ for all the photoanodes. This result shows that charge tunneling in the Bi₂S₃/ NiS and Bi₂S₃/NiS/NiFeO photoanodes occurs primarily through the SS of NiS and the NiFeO co-catalyst (Fig. S10). Although the SS are considered to be hot spots for surface recombination, the SS and catalysts are charged together and reach quasi-equilibrium, resulting in the passivation of the SS. [49] The charge transfer step can be quantitatively described by the interfacial charge transfer rate constant (k_{ct}) derived from R_{ct} . Similarly, the surface recombination can be quantified by the recombination rate constant (k_{rec}) , which is derived from R_{trap} . Both rate constants were calculated using the PEIS data as follows [50, 51]:

$$k_{\rm ct} = \frac{1}{R_{\rm ct}C_{\rm SS/cat}} \tag{2}$$

$$k_{\rm rec} = \frac{1}{R_{\rm trap} C_{\rm SS/cat}} \tag{3}$$

In agreement with the J-V curve, the $k_{\rm rec}$ value of the ${\rm Bi}_2{\rm S}_3$ anode is higher, which results in low net photocurrent flow across the interface. The nonlinear behavior of the $k_{\rm rec}$ values of both the ${\rm Bi}_2{\rm S}_3$ and ${\rm Bi}_2{\rm S}_3/{\rm NiS}$ anodes indicates that band bending is modulated by the time dependened concentration of trapped holes at the SS [52]. The high concentration $N_{\rm ss}$ results in charge storage and modifies the potential drop across the Helmholtz layer. However, after the NiS nanoshell layer was added, $k_{\rm rec}$ was significantly reduced throughout the potential window, especially at higher potential ($\geq 0.8 \ V_{\rm RHE}$), as shown in Fig. 4e. These results are consistent with the J-V curves and $\eta_{\rm bulk}$ plots. Overall, the results suggest that the ${\rm Bi}_2{\rm S}_3/{\rm NiS}$ anode is

severely affected by the SS at the interface, which decrease the benefits of n-p junction formation at lower potentials. Although the continuous forward polarization charges all the SS, further band bending (the absence of Fermi-level pinning) allows for the release of charges into the electrolyte. By contrast, the Bi₂S₃/NiS/NiFeO anode shows a monotonic decrease in $k_{\rm rec}$ with potential, indicating less dependence on the applied potential. This result is consistent with the shift in Nss maximum to lower potentials could charge the SS at lower potentials or the estimated high Nss values originally originates from surface-bounded reaction intermediates. The decrease in k_{rec} from Bi₂S₃ to Bi₂S₃/NiS/NiFeO ultimately increases the carrier lifetime. Quantitative measurement of the charge carrier lifetime can potentially be employed to evaluate the degree of charge separation. TRPL spectroscopy was used to determine the lifetime of excited carriers in bismuth sulfide before and after the formation of the n-p junction. The charge dynamics curves were fitted with a biexponential decay function (i=2),

$$I(t) = \sum_{i} A_{i} e^{\binom{-t}{\tau_{i}}}$$
(4)

where I(t) is the TRPL intensity, A is the amplitude as a normalized percentage, and τ is the lifetime [53]. The fitted results are shown in Fig. 4f; τ_1 represents the non-radiative decay of charge carriers generated in the Bi₂S₃ core through the SS. Upon incorporation of NiS (n-p junction), τ_1 increased drastically due to the alleviation of all SS by the electronic equilibrium attained at the Bi₂S₃/NiS interface [43, 54, 55]. Instead of being trapped by SS and recombined, the charge carriers are separated and accumulated at the valance band of NiS and conduction band of Bi_2S_3 . The reduction in τ_2 represents the successful extraction of charge carriers from NiS due to the creation of additional charge relaxation pathways [56, 57]. A carrier lifetime on the nanosecond scale is generally attributed to a short hole diffusion length (the maximum distance traveled by a carrier without recombination), that is, fast recombination. In real-time operation, the radiative and non-radiative lifetimes of charge carriers will be different or long as the photoelectrode will be connected in a complete electrical circuit with back contact and electrolyte to naturally consume the charge carriers. The carrier lifetime could also vary according to the changes in band edge positions upon polarization [58]. In non-contact TRPL analysis, the charge carriers quickly reach their original positions relative to real-time operating conditions when photon energy stops. However, this analysis gives reliable information about photoelectrodes that belong

to the same sub-groups. The PL spectra of Bi_2S_3 shown in Fig. S11 have two main recombination pathways through the intermediate trap stats at 516.47 and 699.22 nm. The incorporation of NiS and NiFeO drastically reduces the fluorescence intensity by relieving recombination, as shown by the TRPL analysis. By contrast, k_{ct} is higher throughout the potential range compared to that of Bi₂S₃/NiS because NiFeO has a lower stoichiometric Ni²⁺ content and better catalytic properties than NiS (Fig. 4g). The shape of the $k_{\rm ct}$ curve is consistent with that of the J–V curve; briefly, the increase in photocurrent in Bi₂S₃/NiS/NiFeO is larger at lower potentials and becomes almost the same as that of Bi_2S_3/NiS at higher potentials. This phenomenon, a lightintensity-dependent photocurrent, is caused by extreme band bending. The depletion region extends beyond the n-p junction and could extract holes from the VB of bismuth sulfide directly. Similar phenomena also appear in the J-V curve of the Bi₂S₃/NiFeO photoanode (Fig. S12). Furthermore, the carrier lifetime increased from 1.10 to 1.54 ns because the presence of NiFeO sharply decreases k_{rec} . From the obtained PEIS parameters, the overall quantum efficiency of all the photoanodes was calculated as follows:

$$\varphi = \frac{k_{\rm ct}}{k_{\rm rec+}k_{\rm ct}} \tag{5}$$

Figure 4h plots the quantum efficiency versus potential; the curve is in excellent agreement with the J-V curve displayed in Fig. 3a. Specifically, k_{ct} is the dominant factor determining the shape of the J-V curve. As stated above, the enhanced PEC performance results from the formation of a multilayer structure with the necessary components (n-p junction, co-catalyst, and passivation layer) [59, 60].

From the above results, the following working mechanism is proposed for the multilayer Bi₂S₃/NiS/NiFeO/TiO₂ photoanodes for PEC hydrogen generation. The band alignments of the photoanodes under a forward bias in the absence of Fermi-level pinning are shown in Fig. 5a-b [61]. The Fermi level $(E_{\rm F})$ of bare Bi₂S₃ is in equilibrium with the redox potential; both the VB and conduction band (CB) undergo upward band bending within the space-charge region, which is labeled W, and the magnitude of band bending is controlled by the forward bias and the corresponding alignment of $E_{\rm F}$. Under illumination, the photoanode produces an inherent photovoltage (V_{ph}) of 222 mV, which drives the holes into the electrolyte and results in upward shifts of the CB and VB edges, as shown in Fig. 5a [53, 62]. This potential is sufficiently weak to allow effortless charge recombination at a higher rate (k_{rec}) ; as a result, only a fraction of the generated holes tunnel into the electrolyte. When the n-p junction is formed between Bi₂S₃ and the NiS nanolayer, an interior electric field is established because of electron sharing at the junction (Fermi-level equilibrium), as shown in Fig. 5b. The electric field causes spatial charge separation, which causes holes to collect in the VB of NiS and electrons to collect in the CB of Bi₂S₃. However, the existing SS comprise the actual port for charge tunneling, which consumes the photogenerated holes for charging as well as the forward bias. This significantly degrades the utilization of the n-p junction at lower bias, as indicated by the J-Vcurves and η_{bulk} plots. An ultrathin NiFeO co-catalyst layer effectively passivates the intrinsic SS in NiS and accelerates charge tunneling through the NiFeO itself, as shown in Fig. 5c. It drastically reduces the probability of recombination, which increases the carrier lifetime and ultimately the charge injection. Briefly, the multilayer configuration reduces the minimum bias required for charge tunneling by shifting the active SS to lower potential. Finally, the TiO_2 passivation layer on the outside of the electrode prevents the physical degradation of the active layers in the multilayer photoanode. The effects of the TiO₂ passivation layer on the durability and interfacial kinetics are discussed in the next section. This unique thin-film multilayer strategy for efficient electrochemical interface kinetics is potentially useful for many types of electrochemical energy conversion and storage applications.

3.4 Integration of Electrocatalytic and Photoelectrochemical Systems for Noble-metal-free Water Splitting

We verified the durability of the $Bi_2S_3/NiS/NiFeO$ photoanode using chronoamperometry (CA) at a potential of 1.0 V_{RHE} under chopped illumination. The *J*–*T* curve in Fig. 6a shows that the $Bi_2S_3/NiS/NiFeO$ photoanode retained only 58.35% of its initial photocurrent after 1 h of operation due to the degradation of NiS and Bi_2S_3 during prolonged highcurrent–density operation. A thin layer of TiO₂ (~10 nm) was deposited over the $Bi_2S_3/NiS/NiFeO$ photoanode as third functional layer to avoid direct contact with the electrolyte and maintain the structural integrity during highcurrent–density operation. The HR-TEM image in Figs. 6b and S13 shows clear boundaries between the Bi_2S_3 core, NiS nanoshell, NiFeO co-catalyst, and TiO₂ with an approximate thickness of 1 nm represents conformal deposition



Fig. 5 Band alignment of **a** bare Bi_2S_3 , **b** $\text{Bi}_2\text{S}_3/\text{NiS}$, **c** $\text{Bi}_2\text{S}_3/\text{NiFeO}$, and **d** $\text{Bi}_2\text{S}_3/\text{NiS}/\text{NiFeO}/\text{TiO}_2$ under forward bias. At equilibrium, the Fermi levels are aligned by electron sharing, which forms the internal field. Electrons and holes are spatially separated, and the maximum V_{ph} is limited by the positions of the Fermi levels of bismuth sulfide (E_n) and nickel sulfide (E_p). E_{Fn} : electron Fermi level, E_{Fp} : quasi-hole Fermi level

of the NiFeO co-catalyst. The J-V curve of the Bi₂S₃/NiS/ NiFeO photoanode before and after the addition of the TiO₂ buffer layer is shown in Fig. 6c. The small decrease in photocurrent represents increased R_{ct} at the electrode—electrolyte interface. The Nyquist plot in Fig. S14 shows that the higher R_{ct} values originate from TiO₂ passivation layer. However, the Bi₂S₃/NiS/NiFeO/TiO₂ photoanode showed significantly improved durability and retained 89.45% of its initial photocurrent after 1 h of operation.

The Pt electrocathode was replaced with the NiS electrocathode to make an integrated PEC-EC water-splitting cell for noble-metal-free water splitting, as shown in Fig. 6d-e. The individual electrocatalytic activity of NiS in the hydrogen evolution half-cell is shown in Fig. S15. As shown in Fig. 6f, the Bi₂S₃/NiS/NiFeO/TiO₂||NiS PEC-EC cell achieved a current density of 21.2 mA cm⁻² with zero bias (cell voltage, 1.23 V). The *J*–*V* curves of the Bi₂S₃/NiS/ NiFeO/TiO₂ photoanode with Pt and NiS as the electrocathode are shown in Fig. S16. Both electrodes provided identical *J*–*V* curves without an overpotential difference to facilitate water oxidation. Chronopotentiometry studies of the Bi₂S₃/NiS/NiFeO/TiO₂||NiS cell were conducted in an air-tight H cell where both electrodes are separated by a Nafion membrane (Fig. S17a). The *J*-*T* curve shows a slight



Fig. 6 a J-T curves of Bi₂S₃/NiS/NiFeO and Bi₂S₃/NiS/NiFeO/TiO₂ photoanodes under chopped illumination, **b** HR-TEM image of Bi₂S₃/NiS/NiFeO/TiO₂ photoanode, **c** J-V curves before and after addition of TiO₂ buffer layer, **d**, **e** two-electrode PEC-EC cell configuration with Bi₂S₃/NiS/NiFeO/TiO₂ and NiS as photoanode and electrocathode, respectively, **f** J-V curve and **g** J-T curve of PEC-EC cell, **h** comparison of experimental H₂ evolution at a current density of 10 mA cm⁻²

increase in applied potential to maintain a photocurrent density of 10 mA cm⁻² during 6 h of operation, Fig. 6g. The J-T curve of the Bi₂S₃/NiS/NiFeOINiS cell largely deviates from a straight J-T profile owing to the detachment of the Bi₂S₃/NiS/NiFeO photoanode from the FTO substrate as shown in Fig. S17b. To accurately measure the amount of H₂ generated, the experiments were performed by connecting the air-tight H cell directly to a gas chromatography instrument with a N₂ (20 sccm) purge. The average yields of H₂ are shown in Fig. 6h. The H₂ evolution rate was 2.85 and 2.42 μ mol min⁻¹ for the Bi₂S₃/NiS/NiFeO/TiO₂||NiS and Bi₂S₃/NiS/NiFeO||NiS cells, respectively.

Finally, seawater splitting experiments were conducted using the same two-electrode H-cell setup with 0.5 M Na_2SO_4 as a supporting electrolyte (pH ~ 7.5). The seawater was used after simple filtration to remove debris and fine sand particles without any chemical treatment (Gamami Beach, Jeollanam-do, South Korea). The *J*–*V* curve of the seawater photoelectrolyzer is shown in Fig. 7a. A photocurrent of 10.44 mA cm⁻² was achieved at 0 V bias (cell voltage, 1.23 V) for seawater splitting at neutral pH. The photoelectrolysis experiment was conducted continuously for 4 h at a current density of 10 mA cm⁻². The measured cell

voltage under dark and illumination conditions is shown in Fig. 7b. The $V_{\rm ph}$ generated to maintain the operating current density is 463 mV, which is almost double the $V_{\rm ph}$ measured in the three electrode configuration. The cell voltage maintained without any increase during 4 h of photo-electrolysis



Fig. 7 a J-V curve of integrated PEC-EC electrolyzer in seawater electrolyte under chopped illumination, b cell voltage measured under photoelectrolysis condition, c comparison of experimental H₂ and O₂ evolution at a current density of 10 mA cm⁻² and calculated faradaic efficiency, d HR-SEM image and, c XPS spectra of Bi₂S₃/NiS/NiFeO/TiO₂ photoanode after 4 h photo-electrolysis

revealed that the TiO₂ passivation layer served the cause well. The H_2 evolution rate (2.71 µmol min⁻¹) is almost twice that of O_2 (1.22 µmol min⁻¹) in seawater electrolyte, as shown in Fig. 7c. The faradaic efficiencies exceeded 96.8% and 93.4% for H₂ and O₂, respectively, indicating that the selectivity of the photoanode is good even with multiple ions present in the seawater electrolyte. The HR-SEM image of Bi₂S₃/NiS/NiFeO/TiO₂ photoanode after 4 h of photo-electrolysis does not show any structural degradation as shown in Fig. 7d. The direct use of seawater $(pH \sim 8.1)$ leads to the formation of Mg(OH)2 and Ca(OH)2 by reactions with OH⁻ ions generated during electrolysis. Furthermore, the HCO₃⁻ bicarbonate ions may react with OH⁻ ions and then, with Ca²⁺ ions to form insoluble CaCO₃ particles [63]. These particles could be adsorbed on the electrodes and hinder the surface reactions. However, no trace of Mg and Cl was found as shown in Fig. 7e. It shows that the TiO_2 surface is non-porous to allow the binding of excess inorganic chloride (NaCl) in the seawater electrolyte thanks to the ALD process. The $2p_{3/2}$ and $2p_{1/2}$ spin orbitals at 346.30 and 350.12 eV in the high-resolution Ca 2p profile represent the adsorption of Ca derivatives on the OER electrode. The atomic percentage of Ca was not significant, and no increase in cell voltage was observed; therefore, binding of Ca derivatives does not affect the electrode activity. The presence of S^{2-} (160.70 eV) and M–O (528.99 eV) peaks in the corresponding S 2p and O 1 s spectra confirms that the surface reconstruction of NiS and NiFeO to corresponding oxyhydroxides is largely prohibited by the TiO₂ passivation layer [64]. The band alignment between Bi_2S_3 and NiS is well preserved. The M-OH peak (530.35 eV) indicates surface attached species from atmosphere or partial conversion of NiFeO during long-term photo-electrolysis.

4 Conclusion

A novel multilayer strategy for constructing a robust photoelectrode structure for PEC water splitting was reported. Each functional layer was carefully crafted by nanometerscale ALD to precisely control the functionality and maintain the shortest electron transport path. The final multilayer Bi₂S₃/NiS/NiFeO/TiO₂ photoanode exhibited a photocurrent density of 33.3 ± 2.1 mA cm⁻² at 1.23 V_{RHE} under AM 1.5 G illumination. Electrochemical interfacial analyses showed that each functional layer in the multilayer assembly played distinct roles in alleviating specific bottlenecks. Finally, noble-metal-free seawater splitting was demonstrated in an integrated PEC-EC cell using NiS as the electrocathode and a $Bi_2S_3/NiS/NiFeO/TiO_2$ photoanode. The multilayer strategy will expand the options for improving interface-dominant electrochemical processes using revolutionary nanoshell architectures, which will contribute to the rational design and development of electrodes for energy conversion and storage.

Acknowledgements We are grateful to Prof. Hong H. Lee for the valuable and in-depth conversations related to this study. This study was financially supported by the National Research Foundation of Korea (2021R1A2C1012735). This research was also supported by Basic Science Research Capacity Enhancement Project through Korea Basic Science Institute (National Research Facilities and Equipment Center) grant funded by the Ministry of Education.

Funding Open access funding provided by Shanghai Jiao Tong University.

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Supplementary Information The online version contains supplementary material available at https://doi.org/10.1007/s40820-022-00822-8.

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