


Cite this: *Nanoscale Adv.*, 2019, 1, 1460

Enhanced charge separation in g-C₃N₄–BiOI heterostructures for visible light driven photoelectrochemical water splitting†

Kazi M. Alam,^{‡a} Pawan Kumar,^{ID ‡a} Piyush Kar,^a Ujwal K. Thakur,^a Sheng Zeng,^a Kai Cui^b and Karthik Shankar^{ID *a}

Heterojunctions of the low bandgap semiconductor bismuth oxyiodide (BiOI) with bulk multilayered graphitic carbon nitride (g-C₃N₄) and few layered graphitic carbon nitride sheets (g-C₃N₄-S) are synthesized and investigated as an active photoanode material for sunlight driven water splitting. HR-TEM and elemental mapping reveals formation of a unique heterostructure between BiOI platelets and the carbon nitride (g-C₃N₄ and g-C₃N₄-S) network that consisted of dendritic BiOI nanoplates surrounded by g-C₃N₄ sheets. The presence of BiOI in g-C₃N₄-S/BiOI and g-C₃N₄-S/BiOI nanocomposites extends the visible light absorption profile from 500 nm up to 650 nm. Due to excellent charge separation in g-C₃N₄/BiOI and g-C₃N₄-S/BiOI, evident from quenching of the carbon nitride photoluminescence (PL) and a decrease in the PL lifetime, a significant increase in photoelectrochemical performance is observed for both types of g-C₃N₄–BiOI heterojunctions. In comparison to heterojunctions of bulk g-C₃N₄ with BiOI, the nanocomposite consisting of few layered sheets of g-C₃N₄ and BiOI exhibits higher photocurrent density due to lower recombination in few layered sheets. A synergistic trap passivation and charge separation is found to occur in the g-C₃N₄-S/BiOI nanocomposite heterostructure which results in a higher photocurrent and a lower charge transfer resistance.

Received 1st October 2018
Accepted 6th January 2019

DOI: 10.1039/c8na00264a

rsc.li/nanoscale-advances

1. Introduction

The search for renewable and environment-friendly alternative energy sources is among the most focused research activities of the current scientific community. Solar energy is abundant, renewable and sustainable. Hydrogen molecules have the potential to act as a portable chemical fuel where harvested solar energy can be stored. The potential of photoelectrochemical water splitting for high capacity storage of solar energy was recognized from an early era.^{1,2} In a typical photoelectrochemical cell involving an n-type semiconductor active layer, electron–hole pairs are generated in the semiconductor photoanode upon illumination of light. Under an applied bias, photogenerated electrons travel through the semiconductor anode and the external circuit to the counter electrode where they participate in water reduction to evolve hydrogen while photogenerated holes are directly transferred from the

photoanode to the electrolyte to perform water oxidation to evolve oxygen. The valence band maximum of the semiconductor photoanode needs to be below the oxidation potential of the oxygen evolution reaction ($\text{H}_2\text{O} + 2\text{h}^+ \rightarrow 2\text{H}^+ + 1/2\text{O}_2$) for efficient hole transfer to the electrolyte. An ideal semiconductor photoanode needs to possess many desirable attributes such as broadband optical absorption (*i.e.* a low bandgap), a high absorption coefficient for visible photons (to generate a large number of electron–hole pairs per unit volume), a high charge separation efficiency, a low charge transfer resistance, plentiful availability of active sites, good chemical and photocorrosion stability, cost-effectiveness *etc.* Traditional photocatalysts such as TiO₂ and composite TiO₂ nanomaterials have been extensively studied in the past.^{3,4} However, wide bandgaps significantly hinder the visible light absorption capacity of these materials, a fact that has promoted an intense search for finding new earth-abundant, photocatalytic semiconductor systems.

Metal-free, low cost and robust graphitic carbon nitride (g-C₃N₄) has proved to be a highly promising potential photocatalyst for producing hydrogen by water splitting since it has a moderate bandgap (2.6 eV) and suitable band-edge positions. This material is abundant in nature, environment-friendly, and has good thermal, photochemical and chemical stability.^{5–7} This semiconductor consists of tri-s-triazine units positioned in

^aDepartment of Electrical & Computer Engineering, University of Alberta, Edmonton, AB T6G 1H9, Canada. E-mail: kshankar@ualberta.ca; Tel: +1 780-492-1354

^bNanotechnology Research Centre, National Research Council of Canada, Edmonton, Canada T6G 2M9

† Electronic supplementary information (ESI) available. See DOI: 10.1039/c8na00264a

‡ These authors contributed equally.



a two-dimensional graphitic polymer structure.⁸ The photocatalytic performance of this material suffers from poor absorption of red and near-infrared photons (due to bandgap value as well as low oscillator strength), a small specific surface area and a rapid carrier recombination rate.^{6,9} Many attempts have been made in order to enhance the photocatalytic performance of graphitic carbon nitride by synthesizing nanosheets, post-calcination, engineering crystal lattice structure, designing nanostructures and forming heterojunctions with other suitable semiconductors.^{6,9–15} In bulk carbon nitride, inter-sheets hydrogen bonding, and multilayered sheets stacking promotes localized interlayer charge recombination processes which significantly reduce the availability of charge carriers.¹⁶ Bulk carbon nitride is composed of stacked carbon nitride sheets randomly interconnected *via* hydrogen bonding between terminal NH_2 and primary or secondary N's ($\text{NH}_2/\text{C}=\text{N}=\text{C}$).¹⁷ The small effective surface area of stacked sheets and localized charge recombination in hydrogen bonded sheets lead to poor intralayer carrier transport. Bulk carbon nitride can be transformed into few layered or monolayer sheets which increase available surface area and reduce inter-sheets hydrogen bonding which improve charge transport and minimize charge recombination process.^{18,19} Various solvent exfoliation approaches have been used for the transformation of bulk carbon nitride into sheets, however the use of such chemicals is undesirable due to tedious workup, hazardous nature and energy intensive ultrasonication or thermal heating.^{20,21} The addition of N rich precursors such as NH_4Cl which liberates NH_3 and HCl gases at elevated temperature facilitates the transformation of bulk sheets into few layered sheets is appealing due to the avoidance of hazardous chemicals involved in chemical exfoliation. During thermal annealing of dicyandiamide and NH_4Cl mixture at high temperature the released NH_3 and HCl from breakdown of NH_4Cl which blew dicyandiamide derived polymers into numerous large bubbles, which behaves as gas templates and yielded few layered $\text{g-C}_3\text{N}_4$ nanosheets.¹¹ Earth abundant bismuth oxyhalides (BiOX , $\text{X} = \text{Cl}, \text{Br}, \text{I}$) constitute another recently discovered ternary compound semiconductor family that has gathered significant attention due to the small bandgaps of compounds in this family (1.7–1.9 eV) and a layered tetragonal crystal structure.^{22–24} In this family, BiOI has been reported to exhibit the highest photocatalytic activities due to its narrow bandgap.²⁵

Both $\text{g-C}_3\text{N}_4$ and BiOI suffer from a low charge separation efficiency. Formation of a heterojunction with these two systems is a viable approach to prevent the recombination of electron–hole pairs. Enhanced light absorption is another attractive avenue for improving the photocatalytic performance of the relatively high bandgap carbon nitride. The two-dimensional platform structure of graphitic carbon nitride makes it a perfect photoactive supporting matrix for bismuth oxyiodide nanostructures. Recently, a handful of papers evaluated the photocatalytic activities of these composites for the photooxidative degradation of environmentally harmful dyes and for CO_2 reduction experiments.^{26–29} Prior reports that examined low bandgap BiOI -based standalone photoanodes or even heterojunction photoanodes have reported remarkably poor photoelectrochemical

performance not exceeding a few microamps of photocurrent in the potential window for water oxidation.^{30,31} Here in this communication, we reported the enhanced photoelectrochemical water splitting performance of these composite systems using both many layered bulk graphitic carbon nitride ($\text{g-C}_3\text{N}_4$ or CN) and few layered carbon nitride sheets ($\text{g-C}_3\text{N}_4\text{-S}$ or CNS) wherein photocurrent densities approaching approx. 1 mA cm^{-2} under AM1.5 one sun illumination are obtained.

2. Experimental section

2.1 Synthesis of $\text{g-C}_3\text{N}_4$ (CN) and $\text{g-C}_3\text{N}_4\text{-S}$ (CNS)

Bulk $\text{g-C}_3\text{N}_4$ (CN) was synthesized by thermal polycondensation of dicyandiamide at 550°C . In brief, 2 g dicyandiamide was taken in an alumina crucible and covered with a lid. The crucible was heated in a tube furnace with a programmed heating of 5°C min^{-1} up to 400°C ; 2°C up to 500°C , and 1°C min^{-1} up to 550°C followed by a dwell time of 2 h at 550°C . Few layered carbon nitride ($\text{g-C}_3\text{N}_4\text{-S}$ or CNS) sheets were prepared by using dicyandiamide and NH_4Cl as precursors with a slight modification in the literature procedure.¹¹ The precursor NH_4Cl releases gaseous products at elevated temperature to facilitate the separation of sheets due to intercalation of gas bubbles in between the sheets and behaves as a gas template. Briefly, 2 g dicyandiamide and 10 g NH_4Cl were mixed together in a mortar. The mixture was then suspended in 50 mL water followed by heating at 100°C until all the water evaporated. Note: dicyandiamide and NH_4Cl can be directly mixed in water too. The obtained solid was crushed, transferred into an alumina crucible, and heated in a tube furnace using a programmed heating rate of 5°C min^{-1} up to 400°C ; 2°C up to 500°C , and 1°C min^{-1} up to 550°C and holding at 550°C for 2 h.

2.2 Synthesis of pristine BiOI , $\text{g-C}_3\text{N}_4/\text{BiOI}$ and $\text{g-C}_3\text{N}_4\text{-S}/\text{BiOI}$ composites

The composite material synthesis was performed using a hydrothermal method available in the literature^{26,27,29} with slight modifications. $\text{Bi}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}$ and ethanolic KI were used to synthesize BiOI . To prepare BiOI and carbon nitride composite, a slight excess of $\text{Bi}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}$ and $\text{g-C}_3\text{N}_4$ (or $\text{g-C}_3\text{N}_4\text{-S}$) (340 mg) were dissolved in 20 mL ethylene glycol (EG) and ultrasonicated for 60 minutes to obtain a homogeneous solution. Separately, a stoichiometric amount of KI was dissolved in ethanol and ultrasonicated until the solution became transparent. Next, the KI solution was added dropwise in the solution containing $\text{Bi}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}$ and $\text{g-C}_3\text{N}_4$ (or $\text{g-C}_3\text{N}_4\text{-S}$) under vigorous stirring. Then the mixed solution underwent continuous stirring for an additional 30 min and was transferred into a Teflon-lined stainless-steel autoclave and heated for 17 h at 150°C . Pristine BiOI was synthesized using same method without adding $\text{g-C}_3\text{N}_4$ (or $\text{g-C}_3\text{N}_4\text{-S}$).

2.3 Photoelectrochemical measurements

Photoelectrochemical water splitting experiments were performed using a three-electrode set-up (CHI 600D potentiostat).



We prepared pristine BiOI, pristine $g\text{-C}_3\text{N}_4$, pristine $g\text{-C}_3\text{N}_4\text{-S}$, $g\text{-C}_3\text{N}_4/\text{BiOI}$ and $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ composite films on fluorine doped tin oxide (FTO) coated glass. First a very thin (~ 50 nm) compact layer of TiO_2 (ref. 32) was deposited on cleaned (water, ethanol and acetone sonicated) FTO followed by drop coating the composite catalysts onto this compact layer. Prior to the drop coating, the catalyst powders were dissolved in alpha-terpineol (30 mg mL^{-1}) and stirred for 60 min. The alpha-terpineol was used as a binder and later evaporated on a hot plate at 200°C . The as-prepared sample on FTO served as the working electrode (photoanode) and Pt was used as the counter electrode (cathode). An Ag/AgCl glass electrode was used as the reference electrode. The electrodes were immersed in an aqueous $0.1 \text{ M Na}_2\text{SO}_4$ electrolyte. Linear sweep voltammetry from -0.8 V to $+0.8 \text{ V}$ with a scan rate of 10 mV s^{-1} was used to monitor the photocurrent response of the photocatalysts mentioned above. The samples were irradiated with AM1.5 G simulated sunlight from a Class A solar simulator (Newport-Oriel Instruments USA). The illumination intensity measured at the surface of photoanode was 100 mW m^{-2} . Further, to probe the visible light induced photocurrent response, the samples were also illuminated using a 425 wavelength LED with an incident power density of 47.70 mW cm^{-2} at the surface of the photoanode. Electrochemical impedance spectroscopy (EIS) was performed using a three electrode configuration at an applied voltage of -0.5 V vs. Ag/AgCl in $0.1 \text{ M Na}_2\text{SO}_4$, with AC amplitude of 0.005 V at a frequency of 100 kHz . Mott-Schottky plots were collected from impedance-potential measurements in $0.5 \text{ M Na}_2\text{SO}_4$ in the -1.0 to $+1.0 \text{ V}$ voltage range at 1 kHz frequency.

2.4 Characterization

The imaging and elemental analyses of $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ and $g\text{-C}_3\text{N}_4/\text{BiOI}$ composites were performed on a JEOL 2200 FS TEM (transmission electron microscope)/scanning TEM operated at 200 kV . EDX mapping was performed under STEM mode with a nominal probe size of 1 nm . The composite powders were dispersed in ultra-dilute suspensions in methanol followed by deposition on carbon-coated copper TEM grids. The surface topographical images of composite films were obtained using a field emission scanning electron microscope (Zeiss Sigma FESEM) with an accelerating voltage of 5 kV . X-ray diffraction patterns of the five different samples (pristine BiOI, pristine $g\text{-C}_3\text{N}_4\text{-S}$, pristine $g\text{-C}_3\text{N}_4$, $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ and $g\text{-C}_3\text{N}_4/\text{BiOI}$ composites) were recorded using a Bruker D8 X-ray diffractometer with a $\text{CuK}\alpha$, radiation source ($\lambda = 1.5406 \text{ \AA}$) operating at 50 W at room temperature and equipped with a 2-D detector (VANTEC-500). The composites that demonstrated the highest photoelectrochemical performance were $60 \text{ wt\% BiOI}$ with $g\text{-C}_3\text{N}_4\text{-S}$ and $40 \text{ wt\% BiOI}$ with $g\text{-C}_3\text{N}_4$. In the rest of this article, we have denoted these samples as $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ and $g\text{-C}_3\text{N}_4/\text{BiOI}$ respectively. A Perkin Elmer Lambda 1050 UV-Vis-NIR spectrophotometer equipped with an integrating sphere accessory was used to collect the absorption spectra through diffuse reflectance spectroscopic (DRS) measurements. Steady state photoluminescence (PL) measurements were performed with a Varian Cary Eclipse fluorimeter using an excitation

wavelength of 360 nm . Time resolved photoluminescence (TRPL) spectra were recorded using a homemade single photon counting system. Samples were photoexcited by a 405 nm picosecond diode laser (Alphas Lasers GmbH) operated at a frequency of 1 MHz , and the emission was detected by a Becker-Hickl HPM-100-50 PMT interfaced to an SPC-130 pulse counter system. This setup has a response time of $\sim 100 \text{ ps}$. Fourier transform infrared (FTIR) spectra were obtained using an Agilent FTS7000 FTIR Imaging System (diamond ATR/attenuated total reflection) and recorded in transmittance mode in the frequency range of $400\text{--}4000 \text{ cm}^{-1}$. Prior to the collection of spectra, powder samples were deposited on the diamond crystal and moderate nitrogen gas flow was maintained through the ATR assembly. Raman spectra were collected using a Nd:YAG laser (Nicolet Omega XR Raman Microscope) with an excitation wavelength of 785 nm .

3. Results and discussion

3.1 Synthesis, morphology, composition and structural analysis

Bulk carbon nitride ($g\text{-C}_3\text{N}_4$) was synthesized by high temperature (550°C) condensation polymerization of dicyandiamide as a carbon nitride framework forming agent. Carbon nitride few layered sheets ($g\text{-C}_3\text{N}_4\text{-S}$) were prepared by thermal annealing of dicyandiamide and excess of ammonium chloride. Dicyandiamide works as a precursor for the carbon nitride framework while ammonium chloride facilitates the formation of few layered sheets due to evolution of excess gaseous products at elevated temperature; the consequent bubble formation facilitates transformation into few layered $g\text{-C}_3\text{N}_4$ nanosheets.¹¹ *In situ* hydrothermal growth of BiOI nanoplates with carbon nitride structures using $\text{Bi}(\text{NO}_3)_3 \cdot 5\text{H}_2\text{O}$ and equimolar ethanolic KI solution precursors was used for the synthesis of heterostructured nanocomposite of carbon nitride/BiOI (Fig. 1).

FESEM image of $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ displays the rough dendritic nanoplate structure of BiOI scaffolded with $g\text{-C}_3\text{N}_4\text{-S}$ framework (Fig. 2a). The high resolution TEM (HRTEM) image of $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ clearly shows two distinct domains. The dense and crystalline domain corresponds to BiOI while less dense amorphous domain was related to $g\text{-C}_3\text{N}_4\text{-S}$ in $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ heterostructure (Fig. 2b). High magnification HRTEM image displayed lattice fringes of BiOI with interplanar d -spacing of 0.30 and

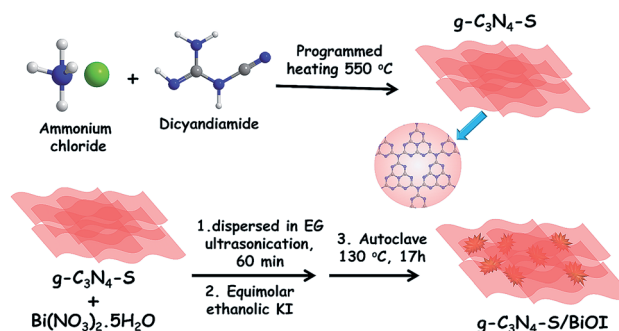


Fig. 1 Synthetic strategy of $g\text{-C}_3\text{N}_4\text{-S}/\text{BiOI}$ heterojunction composite.



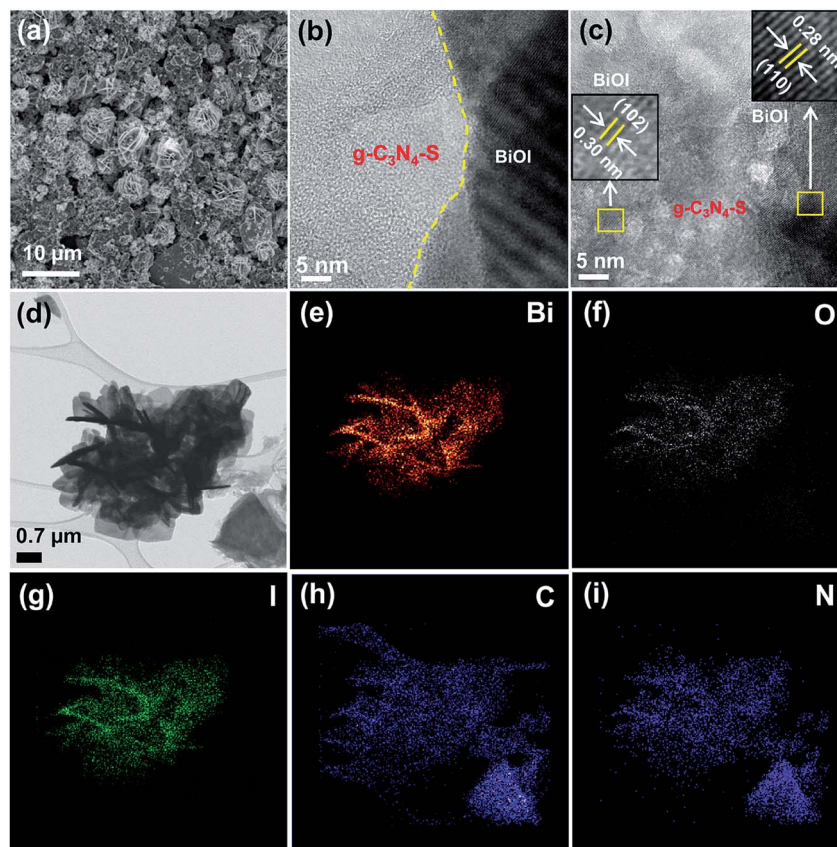


Fig. 2 (a) FESEM image of g-C₃N₄-S/BiOI thin film (b) HRTEM image showing heterojunction of BiOI and g-C₃N₄-S (c) HRTEM image showing lattice fringes (and insets showing *d*-spacing) corresponding to g-C₃N₄-S and BiOI (d) bright field STEM image of g-C₃N₄-S/BiOI; STEM elemental mapping for (e) Bi, (f) O, (g) I, (h) C and (i) N.

0.28 nm assigned to (102) and (110) plane of BiOI respectively (Fig. 2c).^{26,33} Further, various other crystallographic planes with lattice spacings of 0.20, 0.24, and 0.30 nm corresponded to (200), (112), and (102) planes of BiOI respectively were also identified in HRTEM images of g-C₃N₄-S/BiOI.^{26–29} The obtained *d*-spacings were in excellent agreement with XRD results (Fig. 2). Elemental mapping in STEM mode clearly showed even distribution of the constituent elements Bi, O, I, C and N in g-C₃N₄-S/BiOI heterojunction (Fig. 2e–i respectively). The relatively high intensity of C in elemental mapping was due to contribution from carbon of lacy carbon coated TEM grid. The FESEM, HRTEM and elemental mapping images for the bulk g-C₃N₄/BiOI composite system are shown in Fig. 3. g-C₃N₄/BiOI nanocomposite also displayed morphological features identical to g-C₃N₄-S/BiOI except for a more agglomerated sheet structure in g-C₃N₄/BiOI.

Fig. 4a displays the XRD patterns of bulk graphitic carbon nitrides (g-C₃N₄), few layered graphitic carbon nitride sheets (g-C₃N₄-S), bismuth oxyiodide and two heterojunctions (g-C₃N₄/BiOI and g-C₃N₄-S/BiOI). g-C₃N₄ and g-C₃N₄-S show a broad peak centered at ~26° corresponding to the (002) plane originated from interlayer stacking of conjugated aromatic sheets and another small peak at 13.2° corresponding to (100) plane was specific to in-plane repetition of tri-s-triazine unit in the carbon nitride scaffold respectively.⁸ In the composite materials

with BiOI, both (002) and (100) peaks appear, indicative of their preserved crystal structures. The diffractogram of pristine BiOI shows strong peaks characteristic of the tetragonal phase²⁸ and all these peaks appear for both composites, albeit with significantly lower intensities for g-C₃N₄/BiOI compared to g-C₃N₄-S/BiOI. It is worth noting that the BiOI peak for the (001) plane does not appear in g-C₃N₄/BiOI. BiOI has two phases that have small difference. Both phases have a 2D structure along *c* axis and crystallize in the tetragonal space group *P4/nmm*. The difference between the two phases lies in the length of *c* axis (7.21 and 9.14 Å). The phase that shows stronger (001) plane, has shorter *c* axis length.²⁸ The disappearance of the (001) plane in g-C₃N₄/BiOI might be due to (i) the reduction of the BiOI phase that shows (001) plane compared to the other phase due to coupling of (001) plane of BiOI with g-C₃N₄ leading to increased ratio of BiOI with longer *c* axis or (ii) the BiOI nanoplates have higher agglomeration in the g-C₃N₄/BiOI nanocomposite which might be responsible for shielding some signal,²⁸ while in g-C₃N₄-S/BiOI the shielding of signals remain less prominent.

Fig. 4b shows the absorption spectra of the five different samples (pristine BiOI, pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI composites). The large bandgaps of g-C₃N₄-S and g-C₃N₄ significantly limits their response at longer wavelengths, while their heterostructures with BiOI certainly



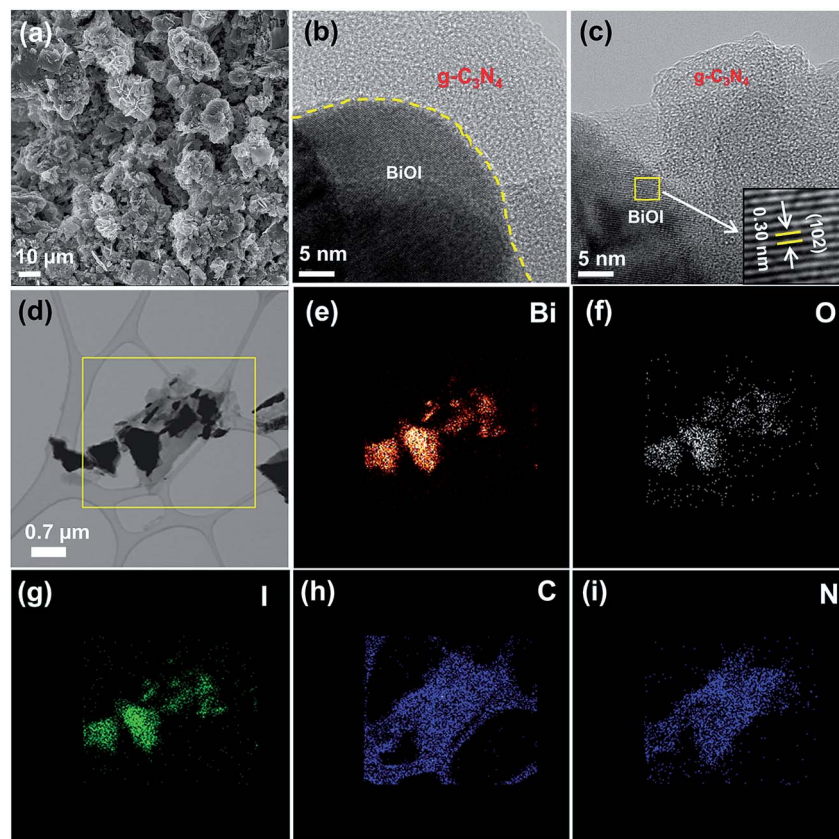


Fig. 3 (a) FESEM image of g-C₃N₄-S/BiOI thin film and HR-TEM of g-C₃N₄-S/BiOI images at (b) 5 nm scale bar showing heterojunction of BiOI and g-C₃N₄-S, (c) at 5 nm scale bar showing lattice fringes and insets shows *d*-spacing corresponding to g-C₃N₄-S and BiOI, (d) bright field STEM image of g-C₃N₄-S/BiOI, and STEM elemental mapping for (e) Bi, (f) O, (g) I, (h) C, (i) N.

have more potential to be a superior photocatalyst. Bulk carbon nitride (g-C₃N₄) shows an absorption band around 270 nm due $\pi \rightarrow \pi^*$ transition while another band around 390 nm with band tailing extended up to 500 nm originates from the $n \rightarrow \pi^*$ transition.³⁴ While both absorption bands were present for few-layered g-C₃N₄-S, the primary band edge was decreased to 450 nm which was attributed to the quantum confinement effect in few layer sheets.²⁰ For the composites, two clear band edges, corresponding to BiOI and carbon nitride, can be seen from the plots. Pristine BiOI exhibits a band-edge at ~ 680 nm with considerable sub-gap absorption extending to 800 nm due to trap states. This sub-gap absorption is suppressed in both g-C₃N₄-S/BiOI and the g-C₃N₄/BiOI, suggesting synergistic trap passivation in the nanocomposite heterostructures. Additionally, bandgaps of the investigated materials were estimated using Tauc plots by extrapolating the linear region of graph between $(\alpha h\nu)^{1/2}$ vs. $h\nu$ on abscissa where α is the absorption coefficient, h is Planck's constant and ν is the light frequency. From the Tauc plots, the values of the electronic bandgap for BiOI and g-C₃N₄ were found to be 1.74 and 2.55 eV respectively while for the few-layered carbon nitride (g-C₃N₄-S), the bandgap was reduced to 2.22 eV due to the formation of sheets (Fig. S4, see ESI†). After hybridization with low bandgap BiOI, the obtained effective bandgap values of the g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructures were found to be 1.79 and 1.84 eV respectively.

Fourier transform infrared spectroscopy (FTIR) was performed to discern the presence of various chemical functionalities in the nanocomposites. The FTIR peaks in 1240 and 1640 cm⁻¹ region for g-C₃N₄-S and g-C₃N₄ were assigned to carbon nitride heterocycle (C–N) stretching modes.³⁵ The other characteristic breathing mode observed at 802 cm⁻¹ originated due to bending vibration of *s*-triazine units (Fig. 4c).^{36,37} BiOI displays a characteristic peak at 521 cm⁻¹ due Bi–O stretch and appearance of this peak in g-C₃N₄-S/BiOI and g-C₃N₄/BiOI composites confirms the presence BiOI in the carbon nitride scaffold.³⁸ Unlike previously reported work,²⁶ we observe a small shift of this peak towards lower wavenumbers which might be due to interaction between BiOI and carbon nitrides.

Fig. 4d shows the Raman spectra for the pristine carbon nitrides (g-C₃N₄-S and g-C₃N₄) and their heterojunctions with BiOI. The two typical bands of carbon-based materials between 1300 and 1600 cm⁻¹ can be found as D and G bands. The out of plane vibrations of sp³ carbon atoms give rise to the D band that represents defects and disorder, and the G band is assigned to E_{2g} phonons which are the in-plane vibrations of sp² carbon atoms characteristics of graphitic structure.^{39,40} Unlike graphene or graphene oxide materials, this bands are not very sharp for relatively amorphous carbon nitrides. The intensity ratio of these bands, I_D/I_G did not change noticeably in the heterostructures compared to the pristine carbon nitrides



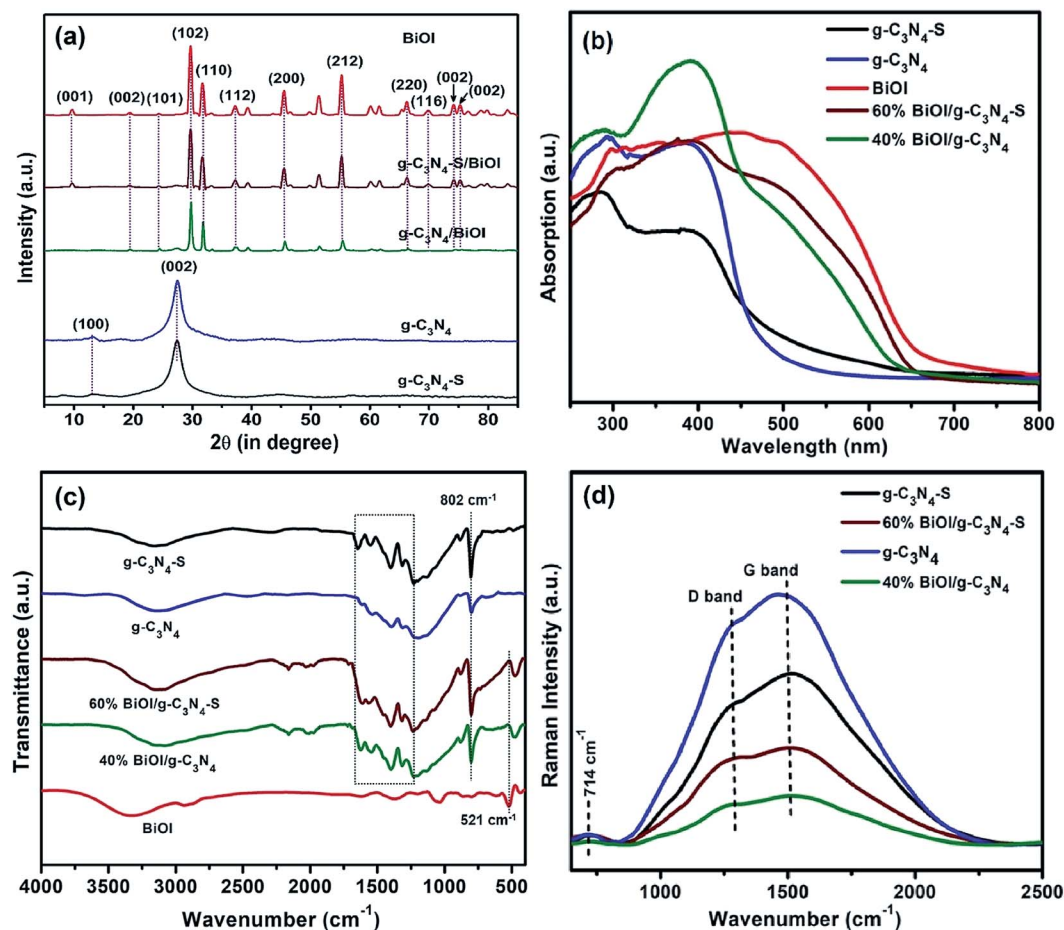


Fig. 4 (a) X-ray diffractograms of pristine BiOI, g-C₃N₄-S/BiOI, g-C₃N₄/BiOI composites, pristine g-C₃N₄ (blue) and pristine g-C₃N₄-S (black) (b) UV-Vis absorption spectra collected in DRS mode (c) FTIR transmission spectra of pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI, g-C₃N₄/BiOI and pristine BiOI (d) Raman spectra ($\lambda_{\text{exc}} = 780$ nm) of pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructures. Color: BiOI (red), g-C₃N₄-S/BiOI (wine red), g-C₃N₄/BiOI (green) and g-C₃N₄-S (black).

(Fig. 7b). This indicates no additional defects have been introduced into the carbon nitride framework in the composites, in other words the in-plane sp² domain size remain unchanged. The small peak around 714 cm⁻¹, which has been attributed to the out of plane C-C vibrations,⁴¹ can be seen as well in the spectrum of both pristine materials and their heterostructures.

3.2 Photocatalytic activity

The photoelectrochemical responses of the samples (pristine BiOI, pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI composites) were obtained under solar simulator (AM 1.5 G (one sun illumination)) using 0.1 M Na₂SO₄ as the electrolyte. Fig. 5a and b show that thin films of both the few layered graphitic carbon nitride sheets (g-C₃N₄-S) and the many layered bulk graphitic carbon nitride (g-C₃N₄) are unable to generate photocurrents higher than 0.20 mA cm⁻² by themselves when used as photoanodes for water splitting. Similarly, the photocurrent density does not exceed 0.45 mA cm⁻² when a pristine BiOI film is used as a stand-alone photoanode (Fig. 5c). However, a photoanode consisting of a nanocomposite film of BiOI and g-C₃N₄-S was able to generate a photocurrent density as high as 0.70 mA

cm² as seen in Fig. 5d. Such a > 50% increase in photoresponse over the pristine, low bandgap BiOI films is indicative of superior charge separation in the nanocomposite thin films. For the g-C₃N₄-S/BiOI composites, the highest photocurrent was observed for 60 wt% BiOI with g-C₃N₄-S and for g-C₃N₄/BiOI composites, 40 wt% BiOI with g-C₃N₄ gave the highest photocurrent. Light on-off experiments (Fig. 5) which demonstrate the photoresponse of the samples, were performed to investigate the charge separation efficiency under illumination. Fig. 5 shows a significant increase in the photocurrent for the heterostructures of carbon nitriles and BiOI compared to their pristine counterparts. These current values are the highest among all values reported thus far for these heterosystems.^{26–29} g-C₃N₄ and BiOI form a Type-II (staggered) heterojunction due to which photogenerated electrons in g-C₃N₄ are transferred to the BiOI while photogenerated holes in BiOI are transferred to g-C₃N₄.^{26,27} This enables both components of the heterojunction to harvest visible light and contribute towards the generation of photocurrent. The ABPE at 0.6 V vs. RHE (water oxidation potential) for BiOI, g-C₃N₄-S, g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI was calculated to be 0.16, 0.05, 0.04, 0.25 and 0.19% respectively (Fig. S8, see ESI†).



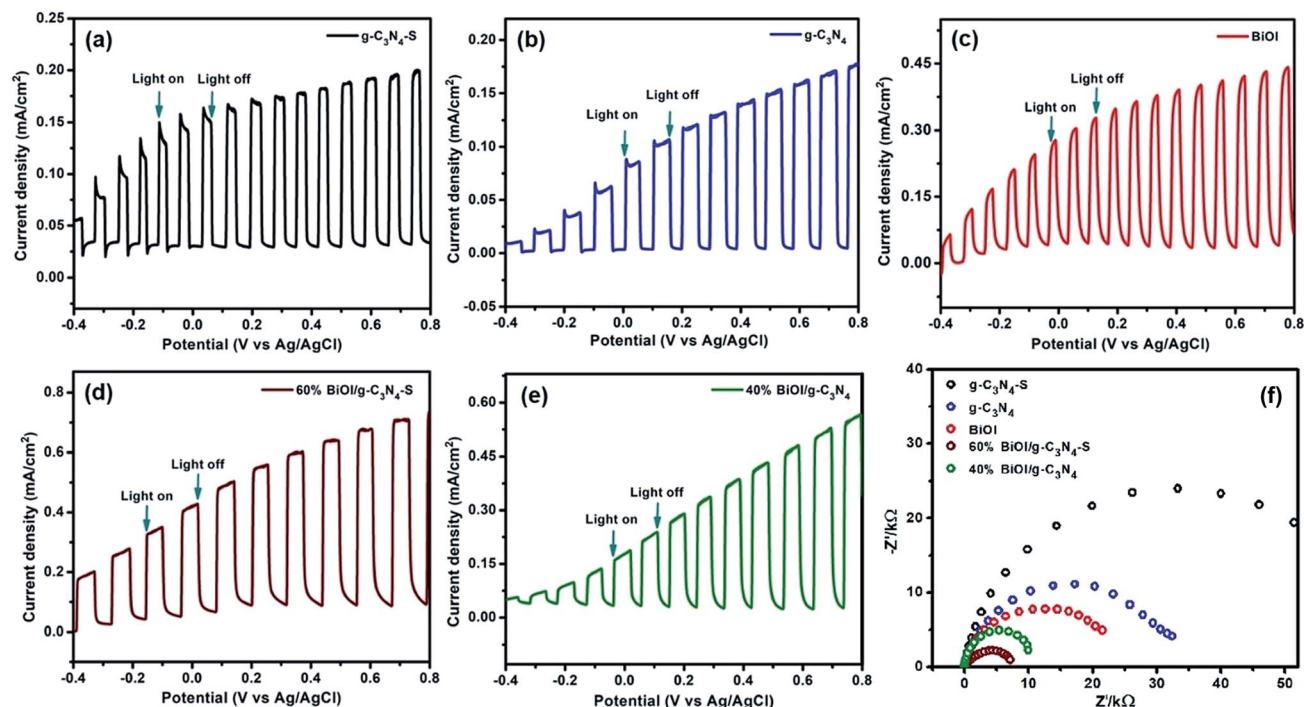


Fig. 5 Light on–off experiment for (a) pristine $g\text{-C}_3\text{N}_4\text{-S}$ (b) pristine $g\text{-C}_3\text{N}_4$ (c) pristine BiOI (d) $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ and (e) $g\text{-C}_3\text{N}_4\text{/BiOI}$; (f) Nyquist plots for the investigated samples obtained from EIS.

Closer examination of the photoresponse curve in Fig. 5a reveals that the photocurrent decreases quickly and monotonically for the pristine $g\text{-C}_3\text{N}_4\text{-S}$ thin film anode for small absolute values of applied bias (up to 0.3 V) whence the electric fields are not strong enough to enforce charge separation – a behavior characteristic of fast carrier recombination. Photocurrent values for the pristine BiOI film (Fig. 5c) do not saturate at a constant value for any value of applied bias during the on cycles and the photocurrent decays are slow during the off cycles, which constitute a signature of extensive trapping of charge carriers. Photocurrent values keep increasing with bias for the $g\text{-C}_3\text{N}_4\text{/BiOI}$ nanocomposite thin film during the illumination on cycles for all values of bias (Fig. 5e), indicating that an intensifying semiconductor–electrolyte potential difference enhances the number of carriers extracted to perform useful work. On the other hand, photocurrent values increase with bias during the on cycles at applied voltages lower than 0.3 V for the $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ nanocomposite thin film (Fig. 5d) and then saturate for higher values of bias, indicating complete extraction of photogenerated charge carriers. The open circuit photovoltage values for pristine $g\text{-C}_3\text{N}_4\text{-S}$ films, pristine $g\text{-C}_3\text{N}_4$ films and pristine BiOI films are -0.58 V, -0.50 V and -0.44 V respectively (vs. Ag/AgCl) while they are -0.56 V in the $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ nanocomposite thin film and -0.64 V in the $g\text{-C}_3\text{N}_4\text{/BiOI}$ nanocomposite thin film. The significant shift in V_{oc} for the $g\text{-C}_3\text{N}_4\text{/BiOI}$ nanocomposite film over the V_{oc} values of the pristine component films confirms increased band-bending and enhanced charge separation at the interface of the electrolyte and the $g\text{-C}_3\text{N}_4\text{/BiOI}$ film while the lack of a similar shift in V_{oc}

for the $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ indicates the quasi-Fermi level of the nanocomposite film to be pinned to that of the $g\text{-C}_3\text{N}_4\text{-S}$ film.

To probe the action of visible photons, we also carried out photoelectrochemical experiments under near-monochromatic 425 nm LED illumination (54.15 W cm^{-2}) under identical conditions. It can be seen from Fig. S2 (see ESI†) that all the materials show a photoresponse for 425 nm irradiation. The highest photocurrent density (0.20 mA cm^{-2}) was observed for the $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ heterostructure while a photocurrent density of 0.16 mA cm^{-2} was found for $g\text{-C}_3\text{N}_4\text{/BiOI}$ which again confirms the symbiotic role of carbon nitride coupled with bismuth oxyiodide in increasing the photocurrent response. For pristine $g\text{-C}_3\text{N}_4$, $g\text{-C}_3\text{N}_4\text{-S}$ and BiOI the value of photocurrent density was found to be 0.05, 0.04 and 0.09 mA cm^{-2} respectively. $g\text{-C}_3\text{N}_4$ (Fig. S2a, see ESI†) and $g\text{-C}_3\text{N}_4\text{-S}$ (Fig. S2b, see ESI†) show transient photoconductivity profiles wherein upon light irradiation, an instantaneous rise in photocurrent was observed, which represents immediate generation of charge carriers and their motion resulting in a photocurrent density spike during the ON cycles. However, this spike suddenly decreased due to fast electron–hole recombination. A steady state photocurrent density was achieved once the rate of electron–hole pair generation and recombination reached a dynamic equilibrium.

For broad spectrum solar illumination, the bulk of the photons are absorbed by the smaller bandgap, more intensely absorbing BiOI component while both the $g\text{-C}_3\text{N}_4$ and BiOI components of the heterostructured nanocomposites absorb equally strongly at 425 nm. As a result, when illuminated by 425 nm photons, the $J\text{-}V$ plots of $g\text{-C}_3\text{N}_4\text{-S/BiOI}$ (Fig. S2d, see



ESI†) and g-C₃N₄/BiOI (Fig. S2e, see ESI†) exhibited subtle but important differences with their *J*–*V* plots under broad spectrum AM1.5 one sun illumination (Fig. 5d and e respectively). During the light ON cycles, photocurrent spikes followed by a rapid decrease in current were seen under 425 nm illumination while the same was not seen under AM1.5 one sun illumination, indicating that recombination was stronger for carriers generated by 425 nm photons. This is because a major portion of the photogenerated minority carriers (holes) in g-C₃N₄ and g-C₃N₄-S experience fast recombination due to a large electron density and concomitant narrow space charge width, as will be shown in Sections 3.3, 3.4 and 3.5. On the other hand, photogenerated electron–hole pairs in BiOI are quickly separated by a strong built-in field, and the low electron density in BiOI means that minority carriers encounter fewer majority carriers to recombine with (see Sections 3.3 and 3.4).

3.3 Electrochemical properties

The semiconductor electrolyte interfacial (SEI) behavior for all the samples is represented by the Nyquist plots (Fig. 5f), which communicate information on the charge transfer resistance (*R*_{ct}). Values of *R*_{ct} can be equated with the diameters of the semicircular arcs denoting the Nyquist plots. As evident from the Nyquist plots, *R*_{ct} decreased in the following order: g-C₃N₄-S > g-C₃N₄ > BiOI > g-C₃N₄/BiOI > g-C₃N₄-S/BiOI. This clearly shows the smaller charge transfer resistance of the nanocomposite heterostructures compared to pristine carbon nitrides. Detailed analysis of the SEI was performed by fitting the Nyquist plots to the equivalent circuit in Fig. S1 (see ESI†), which consists of *R*_{ct} and the Helmholtz capacitance (*C*_h) in the low-frequency (0.1 to 100 Hz) region, a high-frequency (100 to 10 000 Hz) time constant comprising the charge transport resistance (*R*_{sc}) and space charge capacitance (*C*_{sc}), and an electrolyte resistance (*R*_s). The values of circuit parameters, obtained from the equivalent circuit (inset) of the semiconductor–electrolyte interface, are listed in Table 1. Charge transfer processes or diffusion of ions in solution near the SEI are generally slower than electronic processes within the semiconductor.⁴² Therefore, *C*_h and *R*_{ct} were assigned to the low frequency response, and *R*_{sc} and *C*_{sc} were assigned to the high-frequency region. The recombination lifetime ($\tau = R_{sc}C_{sc}$) values, obtained for the SEI, are in the range of several hundreds of milliseconds, and therefore, imply long-lived holes that promote the oxidation of water.^{43–46} Values of τ are listed in Table 1, and are in the following order: g-C₃N₄ > g-C₃N₄-S/BiOI > BiOI > g-C₃N₄/BiOI > g-C₃N₄-S. The very low value of *R*_{ct} = 100 Ω for g-C₃N₄-S/BiOI (Table 1) and the saturation of the

photocurrent at comparatively low bias values for g-C₃N₄-S/BiOI (Fig. 5d) indicates that photogenerated holes in this nanocomposite are able to reach the electrolyte before being captured into trap sites which in turn supports the previously mentioned symbiotic trap passivation effect in g-C₃N₄-S/BiOI.

Charge carrier concentrations and flat band potentials of g-C₃N₄, g-C₃N₄-S, g-C₃N₄/BiOI (40% BiOI/g-C₃N₄), g-C₃N₄-S/BiOI (60% BiOI/g-C₃N₄-S), and BiOI were calculated using the Mott–Schottky relation, given by eqn (1).

$$\frac{1}{C_{sc}^2} = \frac{2}{e\epsilon_0\epsilon_r N_D} \left\{ (V - V_{FB}) - \frac{kT}{e} \right\} \quad (1)$$

$$N_D = \frac{2}{e\epsilon_0\epsilon_r n} \quad (2)$$

In eqn (1), *C*_{sc} is space-charge capacitance per unit area; ϵ_r the dielectric constant of the semiconductor (62 for g-C₃N₄,⁴⁷ and 73 for BiOI^{48,49}); *N*_D is carrier concentration; ϵ_0 is the vacuum permittivity (8.854×10^{-12} F m^{−1}); *k* is Boltzmann constant (1.381×10^{-23} J K^{−1}); *T* is temperature in (298 K); *e* is the electron charge (1.602×10^{-19} C); *V*_{FB} is flat band potential; and *V* is the applied potential. *N*_D, the charge carrier concentration, is calculated from the slope of the Mott–Schottky equations (eqn (1), using eqn (2)). *V*_{FB}, the flat band potential is obtained the point of intersection of the slope (*n* in eqn (2)) with the potential axis as shown in the Mott–Schottky plots (Fig. 6 and S3, see ESI†).

According to Mott–Schottky plot (Fig. 6 and S3, see ESI†), all the semiconductor samples and their heterojunctions investigated in this study are n-type based on the positive slope of the *C*^{−2}–*V* curves. As shown in Fig. S3,† the charge carrier density at equilibrium for g-C₃N₄-S is two orders of magnitude higher than that in g-C₃N₄, highlighting the distinct behavior of few-layer graphitic carbon nitride vs. many-layer bulk carbon nitride. Formation of the heterojunctions with BiOI increases the carrier density by nearly an order of magnitude (over the respective stand-alone photoanodes) for both the g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructure photoanodes, indicating the filling/elimination of electron traps which liberates mobile carriers available for conduction.

3.4 Photoluminescence and lifetime measurement

Fig. 7a shows the steady state photoluminescence (SSPL) spectra of the samples investigated in this work. The g-C₃N₄ and g-C₃N₄-S show intense PL peaks centered at 465 nm. The peak intensity of g-C₃N₄-S was relatively lower than bulk g-C₃N₄

Table 1 Values of *R*_s, *C*_{sc}, *R*_{ct}, *Q* and *n*, obtained by fitting Nyquist plots to the circuit shown in Fig. S2

Sample	<i>R</i> _s (ohms)	<i>C</i> _h (F)	<i>R</i> _{ct} (ohms)	<i>C</i> _{sc} (F)	<i>R</i> _{sc} (ohms)	τ (s)
g-C ₃ N ₄ -S	1	0.000002	42 000	0.000001	16 500	0.0165
g-C ₃ N ₄	1	0.000035	20 000	0.00001	10 000	0.1000
BiOI	1	0.000023	15 000	0.000006	7000	0.0420
g-C ₃ N ₄ -S/BiOI	1	0.000100	100	0.00001	10 000	0.1000
g-C ₃ N ₄ /BiOI	1	0.000090	5000	0.00002	1500	0.0300



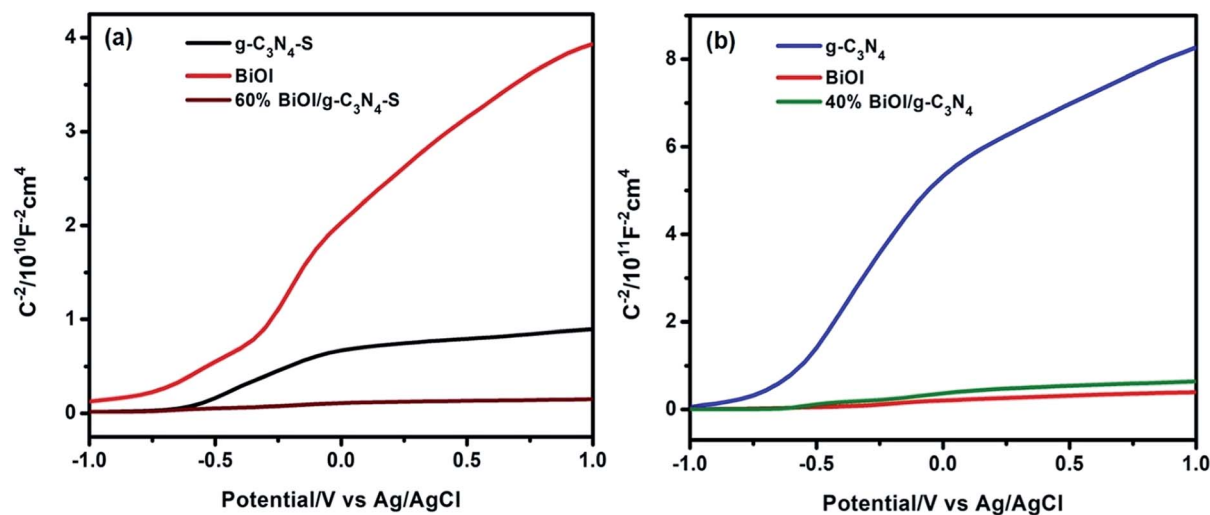


Fig. 6 Mott-Schottky plots for (a) pristine g-C₃N₄-S, pristine BiOI and g-C₃N₄-S/BiOI and (b) pristine g-C₃N₄, pristine BiOI and g-C₃N₄/BiOI. The plots are shown in two different figures with two different scales (vertical axis) for improved visibility.

which represent less prominent charge recombination attributed to better charge carrier separation in few layered sheets. The PL signals for g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructures were quenched significantly compared to the pristine carbon nitrides suggesting efficient charge transfer between BiOI and carbon nitrides. These results were in close agreement with the previously reported works.^{26,27} Small PL peak shift in g-C₃N₄-S/BiOI might originate due to electronic interaction between BiOI and few layered g-C₃N₄-S. To understand the nature of charge recombination processes and determine the lifetimes of excited species, time resolved photoluminescence (TRPL) spectra were measured. The obtained TRPL decay curves of all materials were fitted to a tri-exponential decay function of the form:

$$I(t) = A_1 e^{-t/\tau_1} + A_2 e^{-t/\tau_2} + A_3 e^{-t/\tau_3} \quad (3)$$

where, A_1 , A_2 and A_3 are relative fractional contributions of each decay component and τ_1 , τ_2 and τ_3 are the lifetimes of each

component respectively. Table 2 shows the decay constants, fractional contributions and goodness-of-fit parameters for the pristine carbon nitrides and their heterostructures with BiOI. The presence of three lifetimes in each of the fitted curves is in good agreement with previous reports on carbon nitride-based materials.

For the bulk multi-layered g-C₃N₄, the first and relatively dominant PL decay component has a short lifetime of 2.23 ns, which was attributed due to direct band-to-band radiative recombination (antibonding π^* MO $\rightarrow$ bonding π MO transition). The second PL decay component with intermediate lifetime (12.75 ns) appeared due to radiative decay of electrons from antibonding π^* to LP orbitals *via* intersystem crossing.⁵⁰ The third decay component with a relatively longer lifetime of 144.47 ns likely originated due to inter-sheet recombination processes. The few-layered carbon nitride (g-C₃N₄-S) has two lifetime components (1.97 ns and 12.30 ns), similar in relative intensity and duration to the two shorter lifetime components

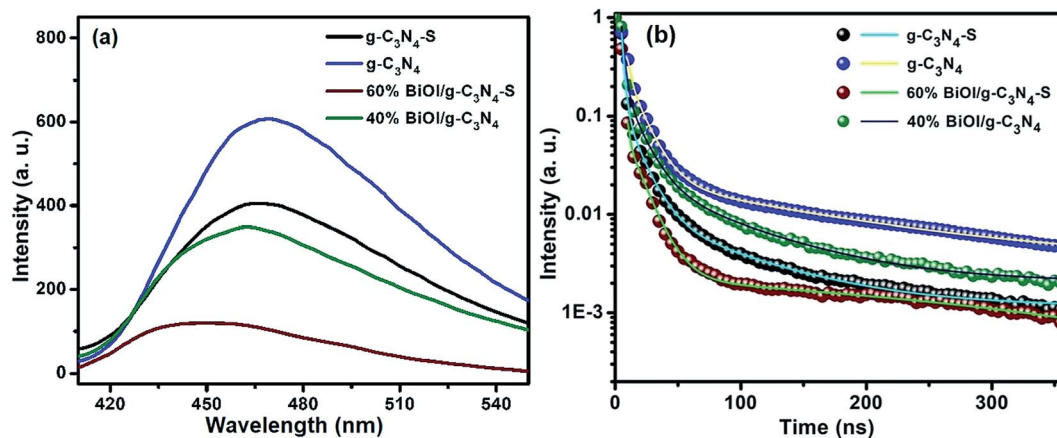


Fig. 7 (a) Steady state PL spectra ($\lambda_{\text{exc}} = 330$ nm) for pristine g-C₃N₄-S, pristine g-C₃N₄, pristine BiOI, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI. The PL spectrum of BiOI was very weak and is not shown. (b) Time resolved photoluminescence spectra ($\lambda_{\text{exc}} = 405$ nm) and (b) Raman spectra ($\lambda_{\text{exc}} = 780$ nm) of pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructures.



Table 2 Summary of fit parameters and goodness of fit information for time-resolved photoluminescence plots shown in Fig. 7a for pristine g-C₃N₄-S, pristine g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI heterostructures

Sample	τ_1 (ns)/ A_1	τ_2 (ns)/ A_2	τ_3 (ns)/ A_3	Average lifetime (τ_{ave}) ns	Red. χ^2	Adj. R^2
g-C ₃ N ₄ -S	74.545/0.0107	12.301/0.176	1.971/7.127	6.98	8.52×10^{-8}	0.999
g-C ₃ N ₄	2.230/11.026	12.746/0.0496	144.472/0.021	16.8	3.24×10^{-8}	0.999
g-C ₃ N ₄ -S/BiOI	3620.125/0.015	12.646/0.119	0.621/279136.0	1.75	1.19×10^{-7}	0.999
g-C ₃ N ₄ /BiOI	12.770/0.268	73.59/0.023	2.213/5.803	10.95	1.33×10^{-7}	0.999

in multi-layered g-C₃N₄ and originates from the same electronic transitions. In contrast to multilayered g-C₃N₄, in few layered g-C₃N₄ sheets, the inter-sheet recombination is reduced resulting in the dramatically reduced relative intensity of the longest lifetime component in g-C₃N₄-S. While the values of the decay components for the g-C₃N₄ and BiOI composites (g-C₃N₄-S/BiOI and g-C₃N₄/BiOI) were changed to significant extents, the relative lifetime decays followed similar patterns to the pristine carbon nitride materials.

The average lifetime (τ_{avg}) was used to evaluate the effectiveness of charge separation in the heterostructures and was determined from the tri-exponential components using the following expression:

$$\tau_{\text{avg}} = (A_1\tau_1^2 + A_2\tau_2^2 + A_3\tau_3^2)/(A_1\tau_1 + A_2\tau_2 + A_3\tau_3) \quad (4)$$

The average lifetimes of g-C₃N₄-S, g-C₃N₄, g-C₃N₄-S/BiOI and g-C₃N₄/BiOI were found to be 6.98, 16.80, 1.75 and 10.95 ns respectively. For BiOI and g-C₃N₄ composites (g-C₃N₄-S/BiOI and g-C₃N₄/BiOI), the average lifetimes are significantly decreased which is attributed to better charge separation in the nanocomposite. The decrease in PL lifetime was consistent with quenching in the steady state PL spectra.

3.5 Mechanism of photoresponse enhancement

Based on bandgap and band potential values, we have proposed a plausible mechanism responsible for the increase in photoelectrochemical water splitting performance. From the Tauc plot the bandgap (E_g) values for g-C₃N₄-S, g-C₃N₄ and BiOI were found to be 2.22, 2.55 and 1.74 eV respectively while the flat band potential values obtained from Mott-Schottky plot were found to be −0.66, −0.67 and −0.59 V vs. Ag/AgCl. The flat band potential can be considered as the position of the conduction band minimum (E_c) if the Fermi level lies just below the conduction band. Therefore, from the value of E_g and E_c , the position of the valence band maximum (E_v) for g-C₃N₄-S, g-C₃N₄ and BiOI were estimated to be +1.56, +1.88 and +1.35 V vs. Ag/AgCl. Usually in water splitting experiments, the standard redox potentials values are expressed on NHE scale so that the E_c positions of g-C₃N₄-S, g-C₃N₄ and BiOI were calculated to be −0.458, −0.474 and −0.391 V while the E_v positions were found to be +1.762, +2.076 and +1.548 V respectively on NHE scale at pH 0. To afford water splitting, the bandgap of material should be higher than 1.23 eV with the position of E_c more negative than 0.00 eV vs. NHE at pH-0 to facilitate reduction of protons (H^+/H_2) while the position of E_v should be more positive than 1.23 eV vs. NHE at pH-0 to achieve

water oxidation (H_2O/O_2). Although the band edge positions of all the semiconductors in this study complied with the basic requirements of water splitting, individual components suffered from faster charge recombination processes. In bulk g-C₃N₄, the charge recombination process remains prevalent due to localized interlayer charge recombination. The transformation of bulk sheets into few layers break hydrogen bonds between sheets and increase crystallinity which resulted into better charge transportation on the surface of sheets. Consequently, heterojunction formation of BiOI with few layered g-C₃N₄-S afforded maximum performance in water splitting experiments. From Mott-Schottky plots (Fig. S4, see ESI†), it is evident that the synthesized BiOI has weaker n-type character than g-C₃N₄-S and its Fermi level lies lower than in g-C₃N₄-S. Heterojunction formation of BiOI with g-C₃N₄-S results in a flow of electrons from the CB of g-C₃N₄-S to the CB of BiOI due to Fermi level difference until equilibrium is reached (Fermi level alignment). This leads to the formation of a Type II (staggered) heterojunction as shown in Fig. 8 where photogenerated electrons migrate from BiOI to g-C₃N₄-S and travel through the external circuit to the platinum cathode to be used for the reduction of protons to hydrogen while holes in the VB of BiOI are used for the oxidation of water (or OH^-). However, the depletion region that would occur as a consequence of the formation of a Type-II heterojunction between BiOI and g-C₃N₄-S would also be expected to reduce the electron density of the heterostructured nanocomposite (g-C₃N₄-S/BiOI) which does not occur; instead Mott-Schottky plots indicate both g-C₃N₄-S/BiOI and g-C₃N₄/BiOI to have an order of magnitude higher carrier density than g-C₃N₄-S and g-C₃N₄ (also supported by the observation in EIS of a lower charge transfer resistance in the composites and the UV-Vis spectral observation of the suppression of sub-gap absorption in the composites), which suggests a synergistic trap filling/trap elimination process in the heterostructured photoanodes that liberates mobile carriers and overwhelms the effect of formation of a Type-II heterojunction. It is demonstrated however that heterojunction formation of BiOI with g-C₃N₄-S promotes better charge separation and increases the visible light photoelectrochemical performance.

3.6 Reusability of the heterojunctions

In order to validate the long-term durability of g-C₃N₄-S/BiOI and g-C₃N₄/BiOI materials, we carried out reuse experiments. For the reuse experiment, FTO deposited material was washed with water and dried at 85 °C and reused for the measurement. The reuse results (Fig. S8, see ESI†) showed a negligible drop in the photocurrent response of heterostructured materials which



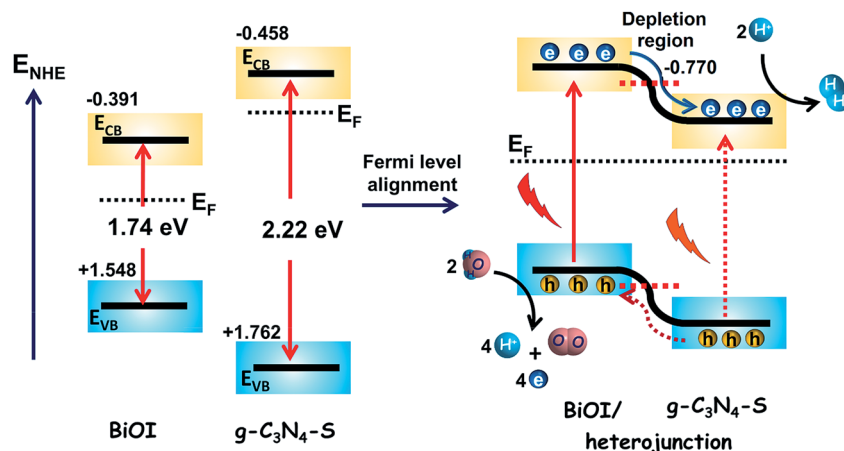


Fig. 8 Proposed mechanism of charge separation in g-C₃N₄-S/BiOI heterostructured photoanodes.

is indicative of longevity and robustness. The used samples g-C₃N₄-S/BiOI and g-C₃N₄/BiOI composites were characterized with diffuse reflectance spectroscopy (DRS) absorption and XRD. Fig. S6 and S7 (see ESI†) show that these data are very similar to the freshly prepared samples. The composite catalysts remain unaffected in their chemical, optical and structural properties after several electrochemical cycles.

4. Conclusion

We have explored the potential of two forms of graphitic carbon nitrides and their heterostructures with earth abundant bismuth oxyiodide. Few layered sheet-like carbon nitride (g-C₃N₄-S) exhibited higher crystallinity, lower inter-sheet recombination and higher carrier densities than multi-layered bulk g-C₃N₄. A significant enhancement in the photocurrent values was observed in the composites under simulated 1 sun illumination with the best performance provided by g-C₃N₄-S/BiOI heterostructured photoanodes. In the heterostructures – more effective charge separation, synergistic trap filling leading to a lower charge transfer resistance, and enhanced light absorption have been identified as dominant factors for the enhancement of photocurrent when compared to the pristine photoanodes. The Mott–Schottky plots revealed both carbon nitrides and bismuth oxyiodide to be n-type semiconductors. Based on the results of experimental characterization and analysis, we identified the mechanism for the electron–hole separation to be the transfer of photogenerated electrons in BiOI to g-C₃N₄-S through the formation of a Type II heterojunction interface. The heterostructured nanocomposites of graphitic carbon nitrides with bismuth oxyiodide displayed almost no loss in photoelectrochemical performance upon re-use and did not suffer damage to their structure and optical properties due to re-use, which bodes well for their long-term operational stability and durability.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank NSERC, NRC, CMC Microsystems, Future Energy Systems and CFI for direct and indirect (equipment use) financial support. U. K. T. acknowledges scholarship support from Alberta Innovates. We acknowledge NRC-NINT and the UofA Nanofab for allowing usage of characterization facilities. Prof. Alkiviathes Meldrum is kindly acknowledged for allowing the use of the TRPL facility in his lab.

References

- 1 K. H. Akira Fujishima, *Nature*, 1972, **238**, 37–38.
- 2 A. J. Nozik, *Annu. Rev. Phys. Chem.*, 1978, **29**, 189–222.
- 3 M. Dahl, Y. Liu and Y. Yin, *Chem. Rev.*, 2014, **114**, 9853–9889.
- 4 P. Kar, Y. Zhang, S. Farsinezhad, A. Mohammadpour, B. D. Wiltshire, H. Sharma and K. Shankar, *Chem. Commun.*, 2015, **51**, 7816–7819.
- 5 A. Naseri, M. Samadi, A. Pourjavadi, A. Z. Moshfegh and S. Ramakrishna, *J. Mater. Chem. A*, 2017, **5**, 23406–23433.
- 6 G. Yue, L. Wang, X. Wang, Y. Chen and D. Peng, *Nanoscale Res. Lett.*, 2009, **4**, 359.
- 7 S. Yin, J. Han, T. Zhou and R. Xu, *Catal. Sci. Technol.*, 2015, **5**, 5048–5061.
- 8 A. Thomas, A. Fischer, F. Goettmann, M. Antonietti, J.-O. Müller, R. Schlögl and J. M. Carlsson, *J. Mater. Chem.*, 2008, **18**, 4893–4908.
- 9 Y. Zheng, L. Lin, B. Wang and X. Wang, *Angew. Chem., Int. Ed.*, 2015, **54**, 12868–12884.
- 10 Q. Lin, L. Li, S. Liang, M. Liu, J. Bi and L. Wu, *Appl. Catal., B*, 2015, **163**, 135–142.
- 11 X. Lu, K. Xu, P. Chen, K. Jia, S. Liu and C. Wu, *J. Mater. Chem. A*, 2014, **2**, 18924–18928.
- 12 M. R. Gholipour, F. Béland and T.-O. Do, *ACS Sustainable Chem. Eng.*, 2016, **5**, 213–220.
- 13 P. Kumar, U. K. Thakur, K. Alam, P. Kar, R. Kisslinger, S. Zeng, S. Patel and K. Shankar, *Carbon*, 2018, **137**, 174–187.
- 14 D. O. Adekoya, M. Tahir and N. A. S. Amin, *J. CO₂ Util.*, 2017, **18**, 261–274.



- 15 B. Tahir, M. Tahir and N. A. S. Amin, *Appl. Surf. Sci.*, 2017, **419**, 875–885.
- 16 K. Schwinghammer, M. B. Mesch, V. Duppel, C. Ziegler, J. r. Senker and B. V. Lotsch, *J. Am. Chem. Soc.*, 2014, **136**, 1730–1733.
- 17 P. Kumar, R. Boukherroub and K. Shankar, *J. Mater. Chem. A*, 2018, **6**, 12876–12931.
- 18 W. Iqbal, B. Qiu, Q. Zhu, M. Xing and J. Zhang, *Appl. Catal., B*, 2018, **232**, 306–313.
- 19 H. Ou, L. Lin, Y. Zheng, P. Yang, Y. Fang and X. Wang, *Adv. Mater.*, 2017, **29**, 1700008.
- 20 S. Yang, Y. Gong, J. Zhang, L. Zhan, L. Ma, Z. Fang, R. Vajtai, X. Wang and P. M. Ajayan, *Adv. Mater.*, 2013, **25**, 2452–2456.
- 21 J. Zhang, Y. Chen and X. Wang, *Energy Environ. Sci.*, 2015, **8**, 3092–3108.
- 22 D. S. Bhachu, S. J. A. Moniz, S. Sathasivam, D. O. Scanlon, A. Walsh, S. M. Bawaked, M. Mokhtar, A. Y. Obaid, I. P. Parkin, J. Tang and C. J. Carmalt, *Chem. Sci.*, 2016, **7**, 4832–4841.
- 23 H. Cheng, B. Huang and Y. Dai, *Nanoscale*, 2014, **6**, 2009–2026.
- 24 M. Guan, C. Xiao, J. Zhang, S. Fan, R. An, Q. Cheng, J. Xie, M. Zhou, B. Ye and Y. Xie, *J. Am. Chem. Soc.*, 2013, **135**, 10411–10417.
- 25 Z. A. Xi Zhang, F. Jia and L. Zhang, *J. Phys. Chem. C*, 2008, **112**, 747–753.
- 26 J. Di, J. Xia, S. Yin, H. Xu, L. Xu, Y. Xu, M. He and H. Li, *J. Mater. Chem. A*, 2014, **2**, 5340–5351.
- 27 S.-Y. Chou, C.-C. Chen, Y.-M. Dai, J.-H. Lin and W. W. Lee, *RSC Adv.*, 2016, **6**, 33478–33491.
- 28 J. C. Wang, H. C. Yao, Z. Y. Fan, L. Zhang, J. S. Wang, S. Q. Zang and Z. J. Li, *ACS Appl. Mater. Interfaces*, 2016, **8**, 3765–3775.
- 29 H. An, B. Lin, C. Xue, X. Yan, Y. Dai, J. Wei and G. Yang, *Chin. J. Catal.*, 2018, **39**, 654–663.
- 30 C. Chang, L. Zhu, S. Wang, X. Chu and L. Yue, *ACS Appl. Mater. Interfaces*, 2014, **6**, 5083–5093.
- 31 L. Xu, P. Yan, H. Li, S. Ling, J. Xia, Q. Xu, J. Qiu and H. Li, *RSC Adv.*, 2017, **7**, 7929–7935.
- 32 U. K. Thakur, A. M. Askar, R. Kisslinger, B. D. Wiltshire, P. Kar and K. Shankar, *Nanotechnology*, 2017, **28**, 274001.
- 33 C. Pan, J. Xu, Y. Wang, D. Li and Y. Zhu, *Adv. Funct. Mater.*, 2012, **22**, 1518–1524.
- 34 A. Kumar, P. Kumar, C. Joshi, S. Ponnada, A. K. Pathak, A. Ali, B. Sreedhar and S. L. Jain, *Green Chem.*, 2016, **18**, 2514–2521.
- 35 S. C. Yan, Z. S. Li and Z. G. Zou, *Langmuir*, 2009, **25**, 10397–10401.
- 36 Y. Wang, R. Shi, J. Lin and Y. Zhu, *Energy Environ. Sci.*, 2011, **4**, 2922–2929.
- 37 J. Zhang, M. Zhang, G. Zhang and X. Wang, *ACS Catal.*, 2012, **2**, 940–948.
- 38 S.-T. Huang, Y.-R. Jiang, S.-Y. Chou, Y.-M. Dai and C.-C. Chen, *J. Mol. Catal. A: Chem.*, 2014, **391**, 105–120.
- 39 A. C. F. a. J. Robertson, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **61**, 14095–14107.
- 40 M. S. Dresselhaus, A. Jorio, M. Hofmann, G. Dresselhaus and R. Saito, *Nano Lett.*, 2010, **10**, 751–758.
- 41 A. K. M. S. Chowdhury, D. C. Cameron and M. S. J. Hashmi, *Thin Solid Films*, 1998, **332**, 62–68.
- 42 T. Lopes, L. Andrade, F. Le Formal, M. Gratzel, K. Sivula and A. Mendes, *Phys. Chem. Chem. Phys.*, 2014, **16**, 16515–16523.
- 43 A. J. Cowan, C. J. Barnett, S. R. Pendlebury, M. Barroso, K. Sivula, M. Grätzel, J. R. Durrant and D. R. Klug, *J. Am. Chem. Soc.*, 2011, **133**, 10134–10140.
- 44 M. Barroso, S. R. Pendlebury, A. J. Cowan and J. R. Durrant, *Chem. Sci.*, 2013, **4**, 2724–2734.
- 45 F. M. Pesci, G. Wang, D. R. Klug, Y. Li and A. J. Cowan, *J. Phys. Chem. C*, 2013, **117**, 25837–25844.
- 46 I. Concina, Z. H. Ibupoto and A. Vomiero, *Adv. Energy Mater.*, 2017, **7**, 1700706.
- 47 S. Pareek and J. K. Quamara, *J. Mater. Sci.*, 2018, **53**, 604–612.
- 48 J. Hu, S. Weng, Z. Zheng, Z. Pei, M. Huang and P. Liu, *J. Hazard. Mater.*, 2014, **264**, 293–302.
- 49 N. T. Hahn, S. Hoang, J. L. Self and C. B. Mullins, *ACS Nano*, 2012, **6**, 7712–7722.
- 50 Y. Zhang, Q. Pan, G. Chai, M. Liang, G. Dong, Q. Zhang and J. Qiu, *Sci. Rep.*, 2013, **3**, 1943.

