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Removal of tetracycline antibiotic activity in water by stable cubic phase barium stannate-perovskite nanoparticles under energy-efficient blue light LED irradiation†

Nehal Ashok Waghchoure, Atul Patel, Laxman G. Raikar, Demi Gandhi, Kampurath Poduvattil Jayadevan * and Halan Prakash *

Photocatalysis is promising for the degradation of antibiotic pollutants in water, which are known to cause the emergence of harmful antibiotic-resistant bacteria (ARB). Importantly, photocatalytic degradation of antibiotic pollutants in water with loss of antibacterial activity in an energy-efficient manner needs to be explored. Here, for the first time, we report degradation of tetracycline (TC), a known antibiotic pollutant in water, with removal of its antibiotic activity by cubic-barium stannate (BaSnO₃, BSO) perovskite photocatalytic nanoparticles under a blue LED light (367 nm) irradiation, (BSO/LED) system. XRD results revealed a single cubic phase of BSO with a crystallite size of 35 nm. FESEM and TEM images showed that BSO had cubic block-like morphology. BET analysis revealed the mesopores nature of BSO with a surface area of $1.1 \,\mathrm{m}^2$ g⁻¹. XPS confirmed the existence of Ba⁺² and Sn⁺⁴ ions. The bandgap energy of BSO was determined to be \sim 3.2 eV. Importantly, photolysis of BSO under blue LED light caused complete degradation of TC in water, with a pseudo-first-order rate constant (kobs) value determined as 0.0824 min⁻¹. Radical scavenging and ESR results revealed O_2 and holes were profoundly generated in the BSO/LED system, causing degradation of TC along with antibiotic activity removal. Electrical energy per order (E_{FO}) was determined as 13.63 kWh per m³ per order, revealing that the BSO/LED system is energy-efficient. Further, BSO was reused four times, highlighting its stability. Degradation byproducts were predicted to be non-toxic by ecological structure activity relationship analysis. Thus, the study discloses an effective photocatalytic degradation of TC along with removal of its antibacterial activity that is needed to prevent hazardous ARB, by a stable non-leadperovskite nano-photocatalyst under an energy-efficient LED source.

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1. Introduction

Antibiotics are used to treat infectious diseases in humans and animals.¹⁻⁴ However, antibiotic waste leads to the contamination of water bodies.³ Water contaminated with antibiotics has been reported to cause the emergence of antibiotic-resistant bacteria (ARBs), which can cause severe disease.⁵⁻⁸ Antibiotics such as tetracycline (TC) are highly consumed.^{9,10} Importantly, TC is not completely metabolised by animals and humans.^{11,12} Thus, TCs are excreted, and their presence in sewage wastewater has been reported, which is a serious problem for the environment and the health of humans and animals.¹³⁻¹⁵ Notably, the presence of TC, even in drinking water sources, has been reported.¹⁶ In the aquatic environment, TC has a long half-life

Energy and Environmental Chemistry Laboratory, Department of Chemistry, Birla Institute of Technology and Science, Pilani, K K Birla Goa Campus, Zuarinagar, Goa, 403726, India. E-mail: halanprakash@goa.bits-pilani.ac.in; jayadev@goa.bits-pilani.ac.in

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(34–329 h),^{17,18} which could lead to accumulation and cause the spreading of ARBs into the environment.¹⁹ Hence, effective degradation of TC in water is urgently needed.

The degradation of antibiotics in water has been reported by various methods such as ozonation,20 UV photolysis,21 Fenton/ photo-Fenton,²² and electrochemical process.²³ Photocatalysis is a promising method for the effective degradation of organic micropollutants.24-26 Photoexcitation of photocatalysts produces excited electron and hole pairs, which are known to generate robust redox reactive species that induce degradation of organic pollutants in water.27-30 Nanocrystalline perovskites are a class of promising photocatalytic materials. 24,31,32 Earlier, mainly leadbased perovskites, having a formula MPbO₃, where M = alkaline earth metals, were reported to show excellent photocatalysis.33 However, lead-based perovskites have a limitation due to the toxicity of the lead.34 Hence, the development of non-lead (Pb) based perovskite photocatalysts has gained significant attention. Barium stannate, BaSnO3 (BSO), is emerging as a nonlead (Pb) based perovskite photocatalyst. 35-37 Earlier, the synthesis of BSO photocatalyst has been reported by different methods like

Paper

combustion technique,38 chemical coprecipitation,39 wet chemicals, 40 sol-gel, 41,42 and hydrothermal, 43 The chemical coprecipitation method has advantages such as low cost, repeatability, high yield, high product purity, and no need of organic solvents.44 Earlier, photocatalytic degradation of antibiotics in water by modified BaSnO3-based photocatalysts, such as tungsten-modified BaSnO₃, heterojunction between BaSnO₃ with photocatalysts like TiO2, copper ferrites and others were reported.,45-47 Tables S4 and S5.† Importantly, the ability of BaSnO₃ nanoparticles to degrade an antibiotic pollutant with the removal of its antibacterial activity after photocatalytic treatment needs to be explored. Moreover, earlier BaSnO3-based photocatalysis were investigated using a conventional xenon lamp.36 Importantly, a xenon lamp emits wide wavelengths of light in the UV and visible ranges, and appropriate filters are needed for photocatalysis.48 Thus, a large amount of light is wasted, and xenon lamp sources consume very high power. Notably, bare BaSnO₃ has poor light absorption above 400 nm.^{49,50} Hence, BaSnO₃ photocatalytic activity with light delivered from energyefficient light sources in the UVA region needs to be explored. 51-53

It is important to note that light-emitting diodes (LEDs) with UVA light are energy efficient and are emerging as an efficient light source. ^{54,55} Currently, UVA-LED (367 nm emission maxima), with a wall plug efficiency of 0.046, is commercially available in the market. The wall-plug efficiency of the LED increases as the UVA radiation maximum increases. ⁵⁶⁻⁵⁸ Importantly, the photo excitation of bare BaSnO₃ (BSO) photocatalysts under blue light LED (UVA) irradiation for the photocatalytic degradation of antibiotics micropollutants in water, with the elimination of antibiotic activity and reuse of BSO photocatalyst, needs to be evaluated.

Therefore, for the first time, we aimed to study the photocatalytic degradation of antibiotic pollutant TC by BSO under blue light LED irradiation (BSO/LED system). A blue LED (367 nm), which closely corresponds to the absorption band of BSO (3.3 eV) was used for irradiation. The BSO obtained by coprecipitation method and characterized by different techniques such as X-ray diffraction spectroscopy (XRD), Field emission scanning electron microscopy (FESEM), transmission electron microscopy (TEM), higher resolution transmission electron microscopy (HR-TEM), Brunauer-Emmett-Teller (BET), selected area electron diffraction (SAED), UV visible spectroscopy, Raman spectroscopy and X-ray photoelectron spectroscopy (XPS). The effect of the initial concentration of TC and BSO photocatalyst, the effect of pH, electron spin resonance (ESR), electrochemical analysis, photocatalytic degradation mechanism, degradation byproducts and ecological structure activity relationships (ECO-SAR) analysis and antibacterial activity before and after BSO/LED system treatment was presented. The reusability of the BSO photocatalyst was performed and ICP-OES analysed the stability of the BSO photocatalyst. The $E_{\rm EO}$ value for the degradation of TC by the BSO/LED system was determined.

2. Experiments

2.1 Materials

The precursor, namely, barium nitrate [Ba(NO₃)₂] (98.5% pure, Sigma-Aldrich), stannic chloride [SnCl₄] (98% pure, Sigma-

Aldrich), sodium hydroxide (NaOH) (Merck), and tetracycline (TCI make) were used in the experiments. The chemicals like methanol, formic acid, and acetonitrile required for HPLC analysis were purchased from Finar Chemicals, and were of HPLC grade. Sodium chloride, sodium bicarbonate, sodium hydrogen phosphate, sulphuric acid, sodium hydroxide, iron(III) nitrate nonahydrate, and humic acid were supplied by Thermo Scientific. The above precursors were directly used without any further purification.

2.2 Synthesis of BSO

The aqueous solution of Ba(NO₃)₂ was prepared by dissolving Ba(NO₃)₂ in 25 ml distilled water. The solution of SnCl₄ was prepared by dissolving SnCl₄ in 10 ml of distilled water. The aqueous solution of SnCl₄ was added dropwise in the above $Ba(NO_3)_2$ solution, stirring for 1 h. After the formation of the homogenous solution, the aqueous solution of NaOH was added to it and stirred for 30 min. This solution was heated at 70 °C for 3 h with constant stirring until the precipitate was formed. The obtained precipitate was transferred to a Petri dish and dried at 80 °C for another 3 h. This dry residue was crushed using a pestle-mortar, transferred in an alumina crucible, heated to 200 °C in a muffle furnace for 2 h, followed by washing with 15 ml of distilled water, dried for 30 minutes at 80 °C, and then heated for two hours at 200 °C. The purpose of the washing stage was to remove the contaminants such as carbonates, nitrates, salt, and chlorine ions. The above process was repeated three times, and powder was calcined and annealed at 500 °C and 800 °C each for 2 h, respectively and allowed to cool at ambient temperature. The powder thus obtained was further used for the analysis.

2.3 Characterization

XRD measurements (Bruker D8) with Cu K α radiation (λ = 1.542 Å) from 20° to 80° with a step size of 0.02 and a step time of 0.6 s. The crystallite size of BSO was calculated to be 35 nm from the Scherrer eqn (1).

$$D = \frac{K\lambda}{\beta \cos \theta} \tag{1}$$

A spectrophotometer (Shimadzu UV-2450) was used to analyse the UV-visible diffuse reflectance spectra (UV-DRS), with BaSO $_4$ as a reference. The measurement range for the spectra was 200–800 nm. The bandgap energy of BSO was calculated from Tauc's eqn (2) and was determined to be 3.2 eV.

$$\alpha h \nu = A(h \nu - E_{g})^{n}. \tag{2}$$

A 532 nm Nd-YAG laser 3.2 mW was used as the laser source for recording Raman spectra (LAB RAM HR Horiba France). The Quanta FEG 250 was used to do FESEM analysis and EDS readings. The voltage for the SEM and EDS experiments was 20 kV. LEICA EM ACE 200 was used for gold sputtering. Transmission Electron Microscopy (TEM) (Fig. 1e) was carried out using FEI Tecnai G2 S-Twin, 200 kV. Thermo Fisher Scientific

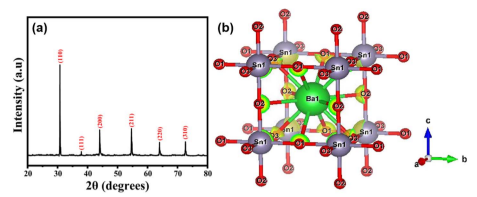


Fig. 1 (a) XRD pattern of single phase BSO synthesised by co-precipitation method, (b) crystal structure of BSO. The centre ball is Ba, and small (red) balls on the edge are the oxygen atoms, and the Sn (violet), which forms an octahedron, is shown coordinating with six oxygen atoms.

(UK) K-Alpha X-ray Photoelectron Spectrometer was used to conduct X-ray photoelectron spectroscopy (XPS).

Electron Spin Resonance (ESR) was recorded using a JEOL ESR Spectrometer. Spin-trapping agents, including DMPO, TEMP, and TEMPO, were used for the analysis. The conditions employed were magnetic centre field = 336 mT, microwave power = 0.995 mW, and microwave frequency of 9.440 GHz. Samples were taken out and examined at the proper intervals.

A BioLogic SP-150e electrochemical workstation was used to carry out the electrochemical analysis. The Mott–Schottky was plotted using electrochemical impedance spectroscopy (EIS) and chronoamperometric measurements, which were performed using a three-electrode electrochemical cell. The Mott–Schottky plots were plotted using EIS measurements, with a reference electrode (Ag/AgCl) having a potential between –2.0 V and –0.4 V with a frequency range of 100 kHz to 10 Hz. The BSO was prepared in ethanol as the working electrode with Nafion acting as a binder. After that, it was applied to a 1 cm² region on fluorine-doped tin oxide (FTO) glass slides. The reference electrode was Ag/AgCl (saturated KCl), while the counter electrode was graphite. For each photoelectrochemical experiment, 15 ml of 0.1 M Na₂SO₄ electrolyte solution, pH = 5.8 and an LED light source (367 nm) were used.

2.4 Photocatalytic degradation experiments

Deionised water was used in the overall photocatalytic experiments. A photocatalytic degradation experiment was conducted in an 80 ml quartz glass reactor at room temperature (25 °C). A desirable amount of TC and BSO were mixed, and the solution was stirred for 30 minutes in the dark. After reaching adsorption equilibrium, 500 μl of the solution was taken in a microcentrifuge tube, which was captioned as 0 min reading and was later centrifuged, and supernatant was used for the analysis. The LED light was switched on and the samples were collected at a specific time interval. An ultraviolet-A lightemitting diode (UVA LED) with a maximum wavelength $(\lambda_{\rm max})$ of 367 nm was used for the degradation experiment. The irradiance spectra of 367 nm LED light is shown in Fig. S5.† The distance between the LED light and the quartz was 0.5 cm, and an external fan was used to keep the LEDs cool. The solution

was stirred continuously throughout the experiment, and 0.5 ml solution was withdrawn at a 3 min time interval in a 1 ml microcentrifuge tube containing 10 μl of methanol as a quencher. This solution was centrifuged at 5000 rpm for 10 min, and the supernatant solution was analysed using high-performance liquid chromatography (HPLC Shimadzu technologies). The Phenomenex Kinetex EVO C18 2.6 μm , 4.6 \times 100 mm column was used for HPLC. An isocratic technique was used with a mobile phase of acetonitrile and formic acid (0.1%) in water (10:90) at a flow rate of 0.8 ml min $^{-1}$. The limit of detection was found to be 0.0944 $\mu g \, L^{-1}$ for TC with an *R*-square value of 0.999 (Fig. S1†).

The following eqn (4) and (5) were used to compute the photocatalytic degradation efficiency (φ) .

$$\varphi = \frac{C_0 - C_t \times 100}{C_0} \tag{3}$$

$$-\ln\left(\frac{C_0}{C_t}\right) = kt\tag{4}$$

To analyse the dynamics data, the first-order model was used:

Where φ is the degradation efficiency, C_0 (mg L⁻¹) and C_t (mg L⁻¹) are the concentrations at the initial and final (after t min), and k is the first order rate constant (min⁻¹). The chemical oxygen demand (COD) was performed to analyse the organic content after the degradation of TC. Initially, the COD before degradation was 9 mg L⁻¹, which was reduced to 4 mg L⁻¹ with an equilibrium time of 45 min for 80 ml TC solution with 0.4 g L⁻¹ of BSO catalyst. The reusability of the BSO catalyst was performed by centrifugation and reuse.

A scavenging experiment was performed using isopropyl alcohol (IPA) as a HO' radical scavenger, sodium azide (NaN₃) as a 1 O₂ scavenger, ethylenediamine tetra acetic acid (EDTA) as a hole (h⁺) scavenger, and 1,4-benzoquinone (BQ) as a superoxide radical anion (O₂ $^{-}$) scavengers. Spin-trapping agents such as TEMP in water and DMPO in water were employed for the analysis of singlet oxygen, and HO', respectively. BSO and DMPO in methanol was used to identify O₂ $^{-}$ radical. ⁵⁹ The aqueous BSO and TEMPO were used to detect holes (h⁺).

2.5 Toxicity assessment

2.5.1 ECOSAR. Acute and chronic ecotoxicity of TC and its intermediate products was predicted by the Ecological Structure–Activity Relationship (ECOSAR application 2.2) developed by USEPA for green algae, fish, and daphnids.^{60–62} The toxicity prediction of organic chemicals is based on available data by structure–activity relationships (SARs). The Globally Harmonized System of Classification and Labelling of Chemicals was used for the categorization of toxicity levels, as shown in Fig. 14.

2.5.2 Antibacterial activity. Antibacterial activity of TC (10 mg L $^{-1}$) before and after treatment by BSO/LED system was assessed by the agar well diffusion method (Clinical and Laboratory Standards Institute, CLSI) as per standard protocol. 63 The antibacterial activity of TC was tested against Gram-positive *Enterococcus faecalis* and Gram-negative *Escherichia coli*. The plates had a final depth of 4 mm after being created in 90 mm sterile Petri dishes with 22 ml of nutrient agar. The spread-plate technique was used to apply 0.1 ml of the tested bacteria's inoculum suspension to the solid media plates. A sterile well borer was used to punched round wells (9 mm in diameter) into agar plates. 100 μ l of samples were filled in the wells. The plates were incubated for 24 h at 37 °C, and the zone of inhibition around the well was measured (mm). Experiments were repeated thrice. 60

2.6 Electrical energy determination

The electrical energy order $(E_{\rm EO})$ was calculated to find the electrical energy required to degrade the contaminant in a water unit. It was calculated as discussed below

$$E_{\rm EO} = \frac{P \times t \times 1000}{V \times 60 \times \log \frac{C_{\rm i}}{C_{\rm f}}} \tag{5}$$

$$\left(\ln\frac{C_{i}}{C_{f}}\right) = k^{'} \times t \tag{6}$$

where P is the power (kW) to drive the LEDs, t is the irradiation time (min), V is the volume (litre), $C_{\rm i}$ and $C_{\rm f}$ are the initial and final concentration of TC and k' is the pseudo-first-order rate constant (min⁻¹) for the decay of TC degradation.⁶⁴ The UVA LED had an electrical input power rating of 2.34 W (12 V and 195 mA, with $E_{\rm max}=367$ nm and FWHM = 15 nm).

From eqn (5) and (6) $E_{\rm EO}$ can be modified into

$$E_{\rm EO} = \frac{38.4 \times P}{V \times k'} \tag{7}$$

Using the above formula, the E_{EO} of the LEDs was calculated as 13.63 kWh per m³ per order.

3. Results and discussion

3.1 Characterization of BSO

3.1.1 XRD analysis. The XRD diffraction peaks centred at 30.68° , 37.80° , 43.93° , 54.53° , 63.87° , and 72.54° corresponds to the planes of $(1\ 1\ 0)$, $(1\ 1\ 1)$, $(2\ 0\ 0)$, $(2\ 1\ 1)$, $(2\ 2\ 0)$, and $(3\ 1\ 0)$ of the cubic phase of BSO (JCPDS 15-0780), (Fig. 1a). The lattice constant for the cubic BSO was calculated as 4.117 Å. The *d*-spacing for the intense peak of the 110 planes at 30.68° was calculated to be 0.2910 nm, which matches the JCPDS file 15-

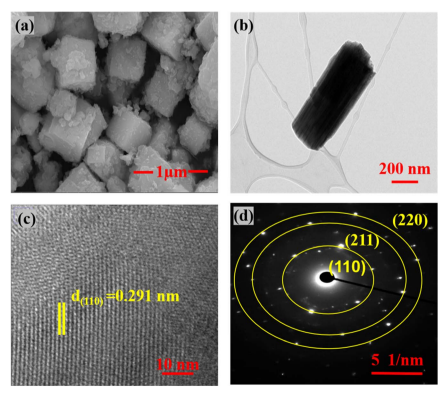


Fig. 2 (a) FE-SEM image of BSO with cubic morphology, (b) TEM image of BSO, (c) HRTEM with d-spacing 0.29 nm, and (d) SAED image of BSO.

0780. The crystal structure of BSO is made using VESTA software⁶⁶ that depicts the positions of all the atoms is shown in Fig. 1b. The crystallite size of BSO was determined to be 35 nm from the Scherrer eqn (1).

3.1.2 FESEM and TEM analysis. FESEM images revealed the block-like morphology of BSO (Fig. 2a). It was observed that the formed cubes are well separated from each other. These cubes are formed via the process of Ostwald ripening. 65 These nanocrystals aggregate to form cubic particles (~1 μm). These cubic particles have rough surfaces. The selected area of the FESEM image and its EDS data are shown in Fig S3a and b,† respectively. A TEM analysis of BSO revealed the nanosheet-like structures that were aggregated to form a block-like morphology (Fig. 2b). The HRTEM image for BSO shows the inter-planar spacing of 0.291 nm, attributed to the lattice plane of 110 (Fig. 2c). This value matches the d-spacing value calculated through the XRD peaks for the 110 planes observed at 30.68°. SAED image of BSO (Fig. 2d) shows the polycrystalline nature of the prepared BSO sample. The bright spots in the SAED image were attributed to the 110 planes corresponding to the most intense peak in XRD.67

3.1.3 Raman, surface area analysis, and absorption property. Two strong bands were observed at 129.2 cm⁻¹ and

147.1 cm $^{-1}$. Importantly, these peaks correspond to the symmetric bending mode (v_5F_{2g}) of [SnO₆] octahedral unit. ^{68,69} Further, the peak at 212.8 cm $^{-1}$ was assigned to the asymmetric bending mode (v_4F_{1u}). ⁷⁰ The bands at 540.3 cm $^{-1}$ and 639.1 cm $^{-1}$ were assigned to the Raman active asymmetric stretching mode (v_2E_g) and symmetrically stretching (v_1A_{1g}) of the Sn–O bond, ⁷¹ respectively (Fig. 3a). The Ba–O bond peak occurred at 408.9 cm $^{-1}$.

The nitrogen adsorption–desorption isotherm of BSO corresponds to a type IV isotherm with an H4 type of hysteresis loop, revealing mesoporous material with narrow-slit pores, as per the IUPAC categorisation of hysteresis loops. The surface area, total pore volume, and pore diameter (BJH) were calculated to be $1.10 \text{ m}^2 \text{ g}^{-1}$, $0.0031 \text{ cm}^3 \text{ g}^{-1}$, and 3.16 nm, respectively, (Fig. 3b and c). The surface area obtained in the present work was slightly lower compared to the earlier suggested report, where BaSnO₃ was obtained *via* microwave-assisted hydrothermal treatment. Importantly, the Tauc plot reveals that BSO has an absorption band edge/tail around 390 nm, and the bandgap energy (E_g) of BSO was calculated to be around 3.2 eV (Fig. 3d).

3.1.4 XPS, Mott-Schottky, and band alignment. Fig. 4a shows the XPS survey of BSO, revealing the presence of Ba, Sn,

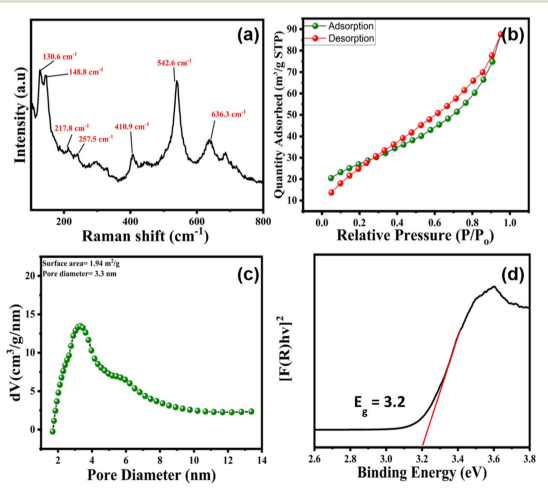


Fig. 3 (a) Raman spectra of pure BSO, (b) nitrogen adsorption—desorption isotherms, (c) BJH pore-size distributions of BSO, and (d) Tauc plot to determine the bandgap of BSO. Minor tick in Fig. 3d.

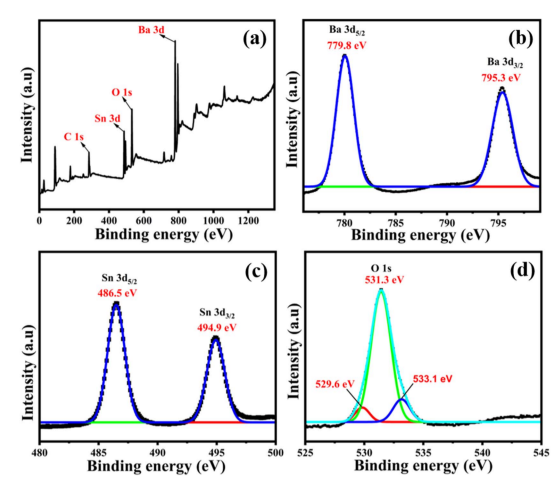
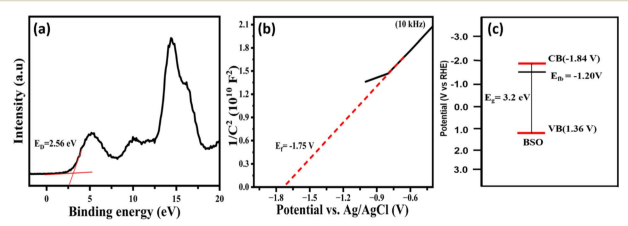


Fig. 4 XPS of bare BSO, (a) XPS survey of BSO, (b) the core-level splitting of peaks of Ba, (c) the core-level splitting of peaks of Sn, and (d) the core-level splitting of peaks of O

and O. The presence of the carbon (C) peak in the XPS survey is attributed to the adventitious carbon from the sample holder.⁷⁷ The Ba 3d XPS spectrum showed two peaks at 779.8 eV and 795.3 eV (Fig. 4b). The difference between two peaks is 15.5 eV, which confirms Ba is having Ba²⁺ oxidation state in BSO.⁷⁸ In Fig. 4c, the Sn 3d XPS spectrum showed two peaks at 486.5 eV and 494.9 eV. The difference between two peaks is 8.4 eV, which confirms Sn is exist in Sn⁴⁺ oxidation state.⁷⁹ O 1s XPS spectrum

showed three deconvoluted O 1s peaks at 529.6 eV, 531.3 eV, and 533.1 eV, which were attributed to lattice oxygen, adsorbed oxygen species in the oxygen-deficient region, and adsorbed hydroxy species, respectively (Fig. 4d). 80,81 These results confirm the presence of Ba, Sn, and O elements in BSO.

Further, the energy difference between the Fermi level and the valence band (E_d) was determined to be about 2.56 V from the XPS valence band spectrum of BSO, Fig. 5a. The Fermi level



(a) XPS valence band spectra, (b) Mott Schottky plot of BSO, (c) band alignment of BSO.

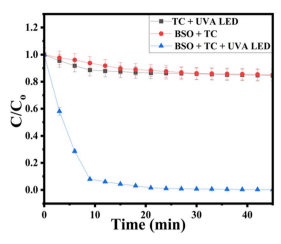


Fig. 6 Degradation of TC by BSO under UVA-LED irradiation. TC = 10 mg L^{-1} , BSO = 0.4 g L^{-1} .

(flat band potential E_f) was determined to be -1.75 V ($\nu s.$ Ag/AgCl) from the Mott–Schottky plot (Fig. 5b), and corresponding E_f value was -1.20 V ($\nu s.$ NHE). Thus, the position of the valence band was determined to be 1.36 V ($\nu s.$ NHE), based on the eqn ($V_b = E_d + E_f$). Further, the position of the conduction band was determined to be -1.84 V ($\nu s.$ NHE) based on the equation $E_{\rm CB} = E_{\rm VB} - E_{\rm g}$, where $E_{\rm g}$ is the band gap energy. The band structure of BSO highlighting the position of VB, CB, and $E_{\rm f}$ is presented in Fig. 5c. A light-emitting diode (LED) that has an emission profile (Fig. S3,† $\lambda_{\rm max} = 367$ nm, 3.3 eV), and strongly overlaps with the absorption (band gap energy of 3.2 eV) of the BSO, was used to evaluate the ability of the BSO photocatalyst (BSO/LED) system to degrade organic pollutant, as discussed below.

3.2 Photocatalysis

3.2.1 Degradation TC by BSO/LED system. BSO/LED photocatalytic system caused 98.42% degradation of **TC** within 45 min as shown in Fig. 6. Notably, the degradation of **TC** by UVA light alone and by BSO alone was not effective in comparison to BSO/LED system. Hence, these results revealed

that adsorption plays no significant role in TC removal, whereas photocatalysis, due to the excitation by the UVA light irradiation, caused the degradation. Excitation of the bandgap of BSO (3.2 eV) with LED causes the generation of electron and hole pairs that are known to initiate the photocatalytic degradation of organic pollutants. Additionally, the degradation of TC under 400 nm LED light with similar irradiance values was also performed and the results showed a reduction in degradation efficiency, with 93.23% degradation achieved in 45 minutes. The higher efficiency observed under 367 nm light (98.42%) is attributed to its better photon energy and stronger absorption by the photocatalyst, leading to enhanced generation of reactive species (Fig. S6†). Further, the effect of BSO concentration, initial concentration of antibiotic pollutant TC, pH effect, and elucidation of the photocatalytic mechanism were investigated, as discussed below.

3.2.2 Effect of concentration of TC on the degradation rate. The rate of degradation decreased with an increase in the initial concentration of TC (Fig. 7a). The pseudo-first-order rate constant values, $k_{\rm obs}$, (Table S3†) were summarised. The $k_{\rm obs}$ values gradually decreased with the increase in the initial concentration of TC (Fig. 7b). This result was attributed to the inner filter effect. The inner filter effect refers to the maximum absorption of light at a relatively higher concentration of TC, which reduces the transmittance of light and prevents the absorption of light and photoexcitation of the BSO. S3,84 Hence, at the comparatively higher concentrations of TC, the reactive radical species involved in the oxidation are not sufficiently available, thereby reducing the degradation efficiency. S5,86

3.2.3 Effect of BSO dosage on the degradation of TC. The TC degradation efficiency was gradually increased with an increase in BSO dosage from 0.1 g L $^{-1}$ to 0.4 g L $^{-1}$, and further increasing the dosage of BSO slightly retarded the rate, exhibiting a bell shape trend Fig. 8b. The results revealed 0.4 g per L dosage had maximum $k_{\rm obs}$ value, and it was considered to be the optimal BSO dosage for the photocatalytic experiments. The effect of BSO catalyst dosage (0.1 g L $^{-1}$ to 0.6 g L $^{-1}$) on tetracycline (TC) degradation was investigated (Fig. 8a).

3.2.4 Effect of pH on the degradation of TC by BSO/LED system. pH of the water is known to influence the surface

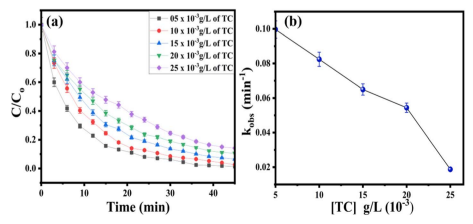


Fig. 7 (a) Effect of TC dosage on degradation rate, and (b) $k_{\rm obs}$ vs. TC g L⁻¹. TC = 10 mg L⁻¹, BSO = 0.4 g L⁻¹.

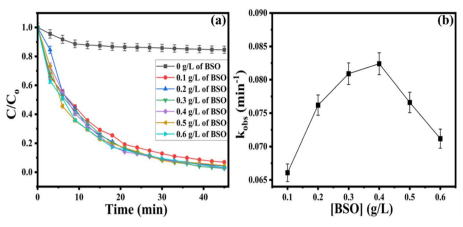


Fig. 8 (a) Effect of BSO dosage on TC degradation, and (b) $k_{\rm obs}$ vs. concentrations of BSO (g L⁻¹). TC = 10 mg L⁻¹, BSO = 0.4 g L⁻¹.

charge of photocatalyst and pollutant, and therefore, the rate of photocatalysis depends on the pH. The $k_{\rm obs}$ values for degradation of TC by BSO/LED system at pH 5 and 6.5 were found to be almost similar, and at pH 9 and 11, there was a slight reduction (Fig. 9a and b). However, at a highly acidic pH 3, there was no significant degradation of TC (Fig. 9a). At a highly acidic pH 3, the zeta potential of BSO is positive, i.e., 9.3 (Fig. S7†). Further, at pH 3 and below, the TC is in protonated form.88 Hence, there is a repulsion between the positively charged photocatalyst, BSO and positively TC molecule at pH 3 (and below), thereby, the degradation is retarded compared to other higher pH of the solution Fig. 9. At pH 5, the zeta potential of BSO is negative. In contrast, the TC molecules are in protonated form, leading to electrostatic attraction between the BSO and TC, and favours photocatalytic degradation compared to pH 3. At alkaline pH 9 and 11, the catalytic surface and the TC molecules acquire negative charges (anionic forms). Thus, catalysts and TC molecules repulsion may cause reduction of the $k_{\rm obs}$ values, in comparison to pH 5 and near neutral pH 7.7.

Interestingly, at highly alkaline pH, the degradation was not significantly reduced compared to acidic conditions at pH 3. This result indicates that the positively charged holes may significantly attract the negatively charged TC molecule under alkaline conditions and cause appreciable degradation, Fig. 9b.

Overall, these results reveal that the BSO/LED system has good degradation ability over a wide pH range.⁸⁹

3.2.5 Effect of inorganic ions and different water matrix on the degradation of TC. Fig. 10a illustrates the effect of tap water and simulated groundwater on TC degradation by BSO/LED system. TC degradation efficiencies in the tap and simulated groundwater were determined to be 63.7% and 46.7%, respectively, compared to 98.42% in Milli-Q water (control), within 45 min. This decline in degradation efficiency is likely due to the presence of inorganic ions and natural organic matter in tap water and simulated groundwater90 (Tables S1 and S2†) that interact with reactive species produced during photocatalysis, thereby hindering TC degradation. Additionally, the BSO/LED system effectively degraded TC at lower initial concentrations (200 μ g L⁻¹ to 1 mg L⁻¹), Fig. S8.† These results suggest that the BSO/LED system is an effective treatment for the degradation of TC pollutants at environmentally relevant concentrations. Earlier, TC pollutant was reported to be in the range of $ng L^{-1}$ to μg L⁻¹ (environmentally relevant concentration) in river and drinking water sources.91

Further, the effect of common inorganic anions in water, *viz.*, Cl⁻, NO₃⁻, HCO₃⁻, SO₄²⁻, and PO₄²⁻ and the effect of organic matter (humic acid, HA) on the degradation of TC by the BSO/LED system was explored. Fig. 10b shows the impact of the

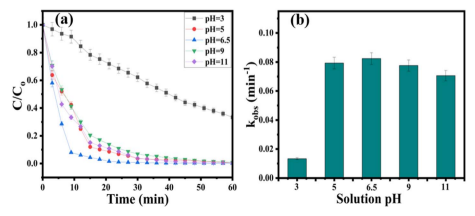


Fig. 9 (a) Effect of initial pH on tetracycline degradation, and (b) $k_{\rm obs}$ vs. pH; [TC] = 10 mg L⁻¹, [BSO] = 0.4 g L⁻¹.

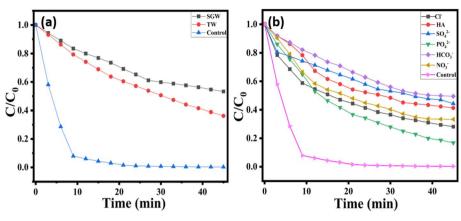


Fig. 10 (a) Effect of water matrices on TC photo-degradation performance by BSO/LED system (b) effects of co-existing inorganic ions and HA and, experimental conditions: [TC] = 10 mg L^{-1} , BSO = 0.4 g L^{-1} , volume: 80 ml, [ion] = 10 mM. SGW = simulated groundwater, TW = Tap water.

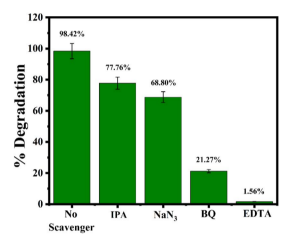


Fig. 11 The effect of scavenging agents on the degradation TC [IPA] = $[NaN_3] = [BQ] = [EDTA] = 100 \text{ mM}$. $[TC] = 10 \text{ mg L}^{-1}$, BSO = 0.4 g L⁻¹.

inorganic ions on the TC degradation efficiency. Among the anions, HCO_3^- exhibits a significant inhibition for the degradation of TC, possibly due to the effective scavenging of HCO_3^- by the h^+ , resulting in decreased photocatalytic activity. Other anions νiz Cl $^-$, NO_3^- and $PO_4^{\,2-}$, may undergo scavenging reactions leading to secondary reactions that produce less reactive species as shown in the reaction eqn (8)–(12) mentioned below. 29,92,93

$$HCO_3^- + h^+ \rightarrow CO_3^{\bullet -}$$
 (8)

$$Cl^- + h^+ \rightarrow Cl^{\bullet}$$
 (9

$$Cl' + Cl^- \rightarrow Cl_2'^- \tag{10}$$

$$NO_3^- + h^+ \rightarrow NO_3^{\bullet} \tag{11}$$

$$HPO_4^{2-} + h^+ \rightarrow HPO_4^{--} + h^+$$
 (12)

Moreover, humic acid (HA), a natural organic compound frequently detected in aquatic environments, significantly impacts the degradation of the tetracycline (TC) BSO/LED system. HA inhibits the degradation process by several mechanisms. Firstly, HA can adsorb onto the photocatalyst's surface, obstructing its active sites and hindering catalytic activity. Additionally, HA exhibits strong absorption in the UVA region, resulting in an inner filter effect that reduces the penetration of UVA light and limits photon availability for photocatalysis. Moreover, HA may react with reactive species generated in the reaction system, reducing the efficiency of TC degradation.⁹⁴

3.2.6 Photocatalytic degradation mechanism. The degradation of TC by the BSO/LED system in the presence of isopropyl alcohol (IPA), sodium azide (NaN₃), benzoquinone (BQ), and EDTA were summarised in Fig. 11. IPA, NaN₃, BQ, and EDTA are known to scavenge HO[•], ¹O₂, O₂ ⁻, and h⁺, respectively. Importantly, EDTA had the highest inhibiting effect compared to other scavengers. Based on these results, the scavenging order was as follows: EDTA > BQ > NaN₃ > IPA. Thus, hole (h⁺) is the predominant species for the degradation of TC, followed by O₂ ⁻, whereas HO[•] and ¹O₂ had a minimum effect. Hence, the proposed BSO/LED system effectively generates reactive species, *viz.*, superoxide and hole for photocatalytic degradation of TC.⁹⁵

ESR was utilized to identify the radicals generated during the photocatalysis. As illustrated in Fig. 12a, the DMPO-superoxide adduct was observed clearly in the BSO/LED system. The oxidation of TEMPO by h⁺, causing a decrease in TEMPO signal, was observed. Further, the photocatalysis reaction in the presence of DMPO showed no noticeable signals of DMPO-OH adduct, revealing HO radical was not effectively generated in BSO photocatalysis Fig. 12b. Additionally, TEMP-singlet oxygen adduct single was not appreciably formed under BSO/LED system. ^{96,97} Fig. 12c. These results corroborate the radical scavenging results, holes, and superoxide radical for the degradation of TC. ^{29,82}

TC under UVA LED irradiation showed no significant degradation (14.2%) because the energy gap between the HOMO and LUMO of TC is 4.2 eV, 98 which is too high for a UVA LED (3.3 eV) to induce direct photochemical excitation and reaction. TC was also not effectively removed by adsorption onto the BSO (Fig. 6). However, 98.42% degradation efficiency was

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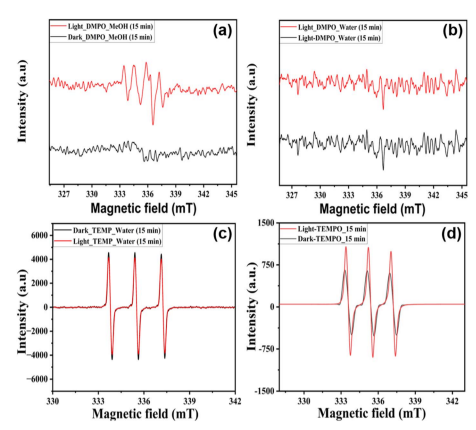


Fig. 12 ESR spectra of (a) DMPO $-O_2^-$ in methanol, (b) DMPO $-HO^+$ in water, (c) TEMP $^-1O_2$ in water, and (d) TEMPO $-h^+$ in water for the dark, and light experiment. ([DMPO] = [TEMP] = 30 mM, BSO = 0.4 g L $^-1$).

achieved by the BSO/LED system within 45 min. Based on the above results, the BSO/LED system photocatalysis mechanism is proposed as discussed below. BSO, upon light irradiation, generates ${\rm e^-}$ and ${\rm h^+}$ pairs. A clear photocurrent response was observed for the BSO under LED irradiation, Fig. S9.† Interestingly, in the presence of TC, the photocurrent response was significantly enhanced, revealing that holes generated in under irradiation were scavenged by the TC. The Nyquist plot (Fig. S10a†) and Bode's plot (Fig. S10b†) were obtained, and the $f_{\rm max}$ (84.64) value was obtained from the Bode plot. The free charge carrier lifetime was determined to be 1.18 ms using eqn (13).⁹⁹

$$\tau = \frac{1}{2\pi f_{\text{max}}} \tag{13}$$

This result further corroborates with the effect of pH, scavenging by the EDTA, and the effect of bicarbonate ion, emphasising that the holes play a pivotal role in degradation next to the superoxide radical ion. The position of the conduction band (CB) -1.84 V (RHE) reveals that BSO is thermodynamically favourable to reducing the oxygen to a superoxide radical anion. Superoxide radical anion is known to induce the oxidation of antibiotic pollutants.²⁹ The $E_{\rm CB}$ value has more negative potential (*i.e.*, -1.84 V) than the standard redox potential of O_2 $-(E(O_2/O_2)^{-1}) = -0.33$ V), which is favourable for

formation of superoxide radicals (O_2^{\cdot}) by the reaction between O_2 and excited photoelectron in CB. ¹⁰⁰ E_{VB} is 1.36 V which is less compared to the standard potential of $HO'(E(HO^{-}/HO')) =$ +1.99 V, and $E(H_2O/HO^*) = +2.73 \text{ V}$, and hence the formation of hydroxy radical (HO') is not favourable, and corroborate well with the ESR, and radical scavenging experiments. 101,102 The position of the valence band (VB) at 1.36 V reveals BSO is not efficient in producing hydroxyl radicals by oxidation of the water molecule. Thus, hydroxyl radicals were not observed in the ESR, and in the presence of hydroxyl radical scavenger, the photocatalysis was not retarded significantly. Under acidic conditions, pH = 3, the O_2 may also go undergo reaction with photogenerated h⁺ to form less reactive H₂O₂ which may also contribute to retard the degradation reaction, in addition to electrostatic repulsion between positively charged TC, and BSO as discussed earlier.103 The degradation mechanism of TC by using BSO/LED light is represented by the following eqn (eqn (14)-(17). 104

$$BaSnO_3 + h\nu \rightarrow e^- + h^+ \tag{14}$$

$$e^- + O_2 \rightarrow O_2^-$$
 (15)

$$h^+ + TC \rightarrow degradation$$
 (16)

$$O_2^{-} + TC \rightarrow degraded TC products$$
 (17)

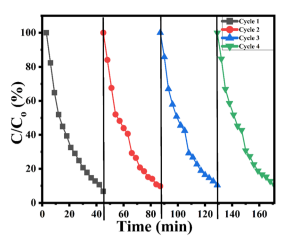


Fig. 13 Recyclability test of BSO photocatalyst, experimental conditions: [TC] = 0.01 g L^{-1} , BSO = 0.4 g L^{-1} , and volume: 80 ml.

3.3 Recyclability and stability studies

The reusability studies of the BSO were studied (Fig. 13). BSO exhibited excellent stability in the reuse study. Importantly, in the fourth cycle, photocatalytic activity was found to be more than 94%. This slight reduction in the activity may be attributed to the loss of the photocatalyst during the washing step. Further, Fig. S2† shows the comparative XRD plots before and

after antibiotic degradation. It was observed that there was no significant decrease in the crystallinity of BSO even after use four times. Additionally, ICP-OES analysis was carried out to evaluate the leaching of active components, and the results showed minimal leaching of Ba (0.18%) and Sn (0.1%), indicating excellent chemical stability. Therefore, the slight reduction in degradation efficiency (from 100% to 94%) is primarily attributed to minor catalyst loss during the washing and recovery steps, rather than any significant deactivation.

The FESEM image shows well-separated smooth-surface cubes even after the degradation experiments (Fig. S3c†). The elemental composition of BSO after TC degradation remains largely unchanged and the atomic percentage and weight percentage of the elements are mentioned in the inset of Fig. S3d.† This indicates the stability of BSO following degradation. After the degradation experiment, the TEM image of BSO was also obtained which revealed no morphological changes (Fig. S4a and b†) compared to the fresh BSO. Overall, these results highlight the stability and reusability of BSO.

3.4 Degradation byproducts and toxicity

Degradation byproducts **TC-1** and **TC-2** were formed due to the cleavage of ring II, with complete ring-opening reactions and bond-breaking reactions, yielding smaller molecular fragments, as shown in Scheme 1. The loss of the side chain amino group in ring IV (deamination) and the amide group in ring IV

Chemical Formula:
$$C_{11}H_{13}NO_6$$

TC-1 (m/z=256)

HO

Chemical Formula: $C_{14}H_{19}NO_2$

TC-2 (m/z=234)

Chemical Formula: $C_{14}H_{19}NO_2$

TC-1 (m/z=445)

HO

Chemical Formula: $C_{22}H_{24}N_2O_8$

TC (m/z=445)

HO

Chemical Formula: $C_{21}H_{21}NO_8$

TC-3 (m/z=404)

Chemical Formula: $C_{21}H_{21}NO_8$

Chemical Formula: $C_{21}H_{21}NO_6$

TC-4 (m/z=384)

Scheme 1 Possible degradation mechanism of TC by BSO/UVA-LED system [TC] = 10 mg L⁻¹, BSO = 0.4 g L⁻¹, and volume: 80 ml.

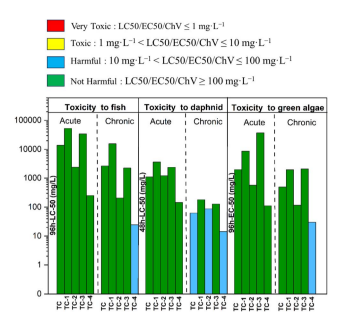


Fig. 14 ECOSAR analysis of TC and degradation byproduct of TC.

(deamidation) led to intermediate TC-3 and TC-4, respectively (Fig. S11 and S12†). Similar degradation byproducts were earlier reported in photocatalysis. 105,106

The ecotoxicity of TC and its degradation by-products using the ECOSAR model were presented in Fig. 14. TC and its degradation byproducts were predicted to be not harmful to acute toxicity towards fish, daphnids (LC50 value > 100 mg L $^{-1}$), and green algae (EC50 value >100 mg L $^{-1}$). Notably, TC and TC-2 showed chronic toxicity against daphnids (categorised as harmful). TC-4 was predicted chronic toxicity towards fish, daphnids, and green algae (categorised as harmful). Importantly, none of the by-products were predicted as being in the toxic chemical category, revealing the BSO/LED system as a potential system for treating TC without generating very toxic degradation byproducts (Fig. 14). 107

3.5 Antibacterial activity assay

The antibacterial activity of TC before and after BSO/LED treatment (at 0–50 min) by BSO photocatalyst was determined (Fig. 15). These results reveal that TC (0.01 g L $^{-1}$) before treatment showed a zone of inhibition against target *E. faecalis* and *E. coli* (Fig. 15A and C). Notably, after treatment (50 min), degraded TC showed no zone of inhibition against *E. faecalis* and *E. coli*. Notably, BSO photocatalysts showed no antibacterial activity against *E. faecalis* and *E. coli* (Fig. 15A and C). These results reveal that treatment by the BSO/LED system caused the complete is first to demonstrate the degradation of TC by BSO/

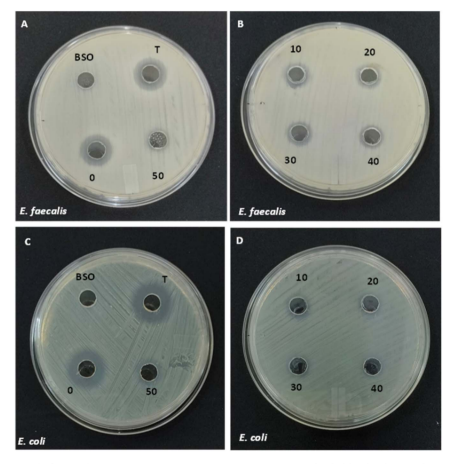


Fig. 15 Anti-bacterial activity of TC and of the treated solution withdrawn at time interval 0, 10, 20, 30, 40, and 50 min against *E. faecalis*, (A and B). Anti-bacterial activity of TC and of the treated solution withdrawn at time intervals 0, 10, 20, 30, 40, and 50 min against *E. coli* (C and D).

LED system with complete removal of the antibacterial activity of TC.¹⁰⁸ To the best of our knowledge, the present study of antibacterial activity is important to prevent the emergence of ARBs.

4. Conclusion

TC was effectively degraded by the proposed BSO/LED system. XRD revealed the cubic phase of BSO. FESEM revealed the cubic morphology of BSO. TEM image showed the aggregation of nanosheets of BSO to form a cubic block-like morphology. The BET analysis reveals the slit-like mesopores in BSO. XPS analysis confirmed the presence of Ba2+, Sn4+ and lattice oxygen and adsorbed hydroxy species in BSO. At the optimised condition (BSO = 0.4 g L^{-1} , TC = 10 mg L^{-1}), the pseudo-first-order rate constant for TC degradation by the BSO/LED system was determined to be 0.0824 min⁻¹. The effects of the initial concentration of TC, dosage of BSO, and pH on TC degradation by the BSO/LED system were presented. The positive Mott-Schottky curve reveals that BSO is an n-type semiconductor with a flat band potential ($E_{\rm fb}$) of -1.75 V (Vs Ag/AgCl). The radical scavenging results highlight the major role of superoxide (O₂⁻) and holes (h+) in the degradation of TC. ESR results confirmed the O₂ and h were present in the BSO/LED system. The photocurrent experiments revealed the interaction of holes (h⁺) with TC, thereby increasing the photocurrent compared to BSO. The antibacterial activity of TC against Gram-negative Escherichia coli (E. coli) and Gram-positive Enterococcus faecalis (E. faecalis) was completely lost after BSO/LED treatment. The degradation pathways involved ring opening, deamination and deamidation reactions to produce degradation byproducts, which were identified by high-resolution mass spectrometry. Importantly, ECOSAR analysis predicted these degradation byproducts to be non-toxic. Further, EEo values were determined to be 13.63 kWh per m³ per order, and the cost was estimated to be \$ 1 m⁻³, revealing that the BSO/LED system is energy-efficient and cost-effective in removing TC. Overall, the finding presents the degradation of TC by the proposed stable and energy-efficient BSO/LED system with antibacterial activity removal which is important to prevent the spreading of ARBs.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There is no conflict to declare.

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