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Visible-light driven p-n heterojunction formed between α -Bi₂O₃ and Bi₂O₂CO₃ for efficient photocatalytic degradation of tetracycline⁺

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To improve the efficiency of photocatalytic oxidative degradation of antibiotic pollutants, it is essential to develop an efficient and stable photocatalyst. In this study, a polymer-assisted facile synthesis strategy is proposed for the polymorph-controlled α -Bi₂O₃/Bi₂O₂CO₃ heterojunction retained at elevated calcination temperatures. The p-n heterojunction can effectively separate and migrate electron-hole pairs, which improves visible-light-driven photocatalytic degradation from tetracycline (TC). The BO-400@PAN-140 photocatalyst achieves the highest pollutant removal efficiency of 98.21% for photocatalytic tetracycline degradation in 1 h (λ > 420 nm), and the degradation efficiency was maintained above 95% after 5 cycles. The morphology, crystal structure, and chemical state of the composites were analysed by scanning electron microscopy, X-ray diffraction, and X-ray photoelectron spectroscopy. Ultraviolet-visible diffuse reflection, transient photocurrent response, and electrochemical impedance spectroscopy were adopted to identify the charge transfer and separation efficiency of photogenerated electron-hole pairs. The EPR results verified h⁺ and 'OH radicals as the primary active species in the photocatalytic oxidation reactions. This observation was also consistent with the results of radical trapping experiments. In addition, the key intermediate products of the photocatalytic degradation of TC over BO-400@PAN-140 were identified via high-performance liquid chromatography-mass spectrometry, which is compatible with two possible photocatalytic reaction pathways. This work provides instructive guidelines for designing heterojunction photocatalysts via a polymer-assisted semiconductor crystallographic transition pathway for TC degradation into cleaner production.

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Introduction

In recent years, with the development of industrialization in the world, wastewater pollution is becoming more and more

serious. The type of pollutants in the water also exhibit a trend of diversification.¹ Emerging antibiotic pollutants, which are considered hazardous to the ecosystem, are prevalent in surface water, groundwater, and wastewater treatment plant effluents.² Among them, tetracycline has often been reported as the most famous broad-spectrum antibacterial agent and antibiotic pollutants in water.³ However, the degradation of tetracycline in wastewater is difficult. Conventional wastewater treatment technologies (chlorine, ozone, filtration, and adsorption) have the disadvantages of high cost, complex processes, limited purification effect, and secondary pollution.⁴ Therefore, it is imperative to develop technologies to effectively reduce the elimination of such antibiotics.⁵

Photocatalysis has attracted significant interest owing to its non-pollution, safety, and other characteristics for addressing environmental pollution.^{6–8} It can degrade organic pollutants in water into carbon dioxide and water. In addition, the photocatalyst used is non-toxic, low-cost, stable and reusable.⁹ Up to now, efforts have been made to develop photocatalysts that use visible light to efficiently remove tetracycline.¹⁰

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Bismuth-based photocatalysts (such as Bi₂O₃, Bi₂O₂CO₃, and Bi_2WO_6) are considered highly favourable candidates for environmental remediation applications owing to their relatively high visible light absorption capacities.¹¹ Notably, α-Bi₂O₃ exhibits excellent photodegradation activity (owing to its favourable bandgap energy structure and efficient photoelectric transformation rate) toward organic pollutants, including Rhodamine B (RhB),¹² 2,4-DCP,¹³ tetracycline (TC).¹⁴ The unique layered structure of n-type Bi₂O₂CO₃ consists of alternating perovskite-like slabs of CO_3^{2-} and layers of $(Bi_2O_2)^{2+}$ to promote photogeneration while also exploiting a limited area of the UV light region.14-16 However, the inefficient utilisation of sunlight and the fast combination rate of photogeneration significantly constrain the degradation activity of Bi2O2CO317 and the construction of a heterojunction is crucial to impeding the recombination rate of photogenerated electron-hole pairs and enhance photocatalytic performance.

In contrast with pure n-type photocatalysts, the generation of an interior electric field between the n-type and p-type semiconductors of the p-n heterojunction effectively drives the photogenerated charge transfer and separation.¹⁸ As a result, the rational bandgap structure of the p-n heterojunction photocatalysts increases the lifetime of the carrier while also preserving the threshold energy level of the photoelectron. Various studies have reported the heterojunction of Bi-based materials, such as Bi₂O₂CO₃/g-C₃N₄ heterojunctions for sulfamethazine degradation¹⁹ and Bi₂O₃/g-C₃N₄ nanocomposites for amido black 10B degradation. Relative to β -Bi₂O₃, α -Bi₂O₃ possesses the advantages of a smaller bandgap, higher oxidative ability of the valence hole, and non-toxicity.20,21 Traditional methods to prepare Bi2O3/Bi2O2CO3 heterojunction photocatalysts are based on heat treatment strategies. In the calcination process, when the calcination temperature reaches about 350 °C, Bi₂O₂CO₃ firstly decomposes to form β-Bi₂O₃, and then transforms from β -Bi₂O₃ to α -Bi₂O₃ by continuing to increase the calcination temperature. At present, the main research is the formation of β -Bi₂O₃/Bi₂O₂CO₃ and β -Bi₂O₃/ α -Bi₂O₃ heterostructure, α-Bi₂O₃/Bi₂O₂CO₃ heterostructure is relatively difficult to achieve. Which are relatively difficult to implement for the successful formation of the Bi2O2CO3/a- Bi_2O_3 heterostructure. For this reason, the use of α - $Bi_2O_3/$ Bi₂O₂CO₃ heterojunction materials in photocatalytic degradation of pollutants has rarely been reported.

Researchers have recently succeeded in implementing the controlled synthesis of photocatalysts through the introduction of polymers as structure-directing agents to control the type of crystal growth.^{22,23} Various polymers with specific functional groups have been utilised owing to their superior ability to anchor with metal ions.^{24,25} Polyacrylonitrile (PAN) has been widely utilised for such controllable syntheses.²⁶ The introduction of small quantities of polymer to the system acts as a protection for partial nanoparticles by enabling them to retain their original crystalline morphology at elevated temperatures.

In this paper, a simple polymer-assisted synthesis method is developed based on the introduction of PAN, thereby enabling the formation of α -Bi₂O₃/Bi₂O₂CO₃ heterostructure directly obtained from Bi₂O₂CO₃ through partial transformation. The $Bi_2O_2CO_3$ precursor was synthesised through the hydrothermal method. The polymer, PAN, was specified as the capping agent to envelop the partial $Bi_2O_2CO_3$ precursor. In addition, TC and RhB were adopted as the representative target pollutants to assess the photocatalytic properties of the α -Bi₂O₃/Bi₂O₂CO₃ heterostructure photocatalyst. This work provides deep insight and a promising strategy to design highly efficient hybrid photocatalysts for environmental remediation.

Experimental section

Chemicals

Ethylenediaminetetraacetate dihydrate (EDTA), bismuth nitrate pentahydrate (Bi(NO₃)₃· 5H₂O), citric acid (C₆O₇H₈), TBA, polyacrylonitrile (PAN, average molecular weight 150 000), and Lascorbic acid (L-AA) were purchased from Aladdin Reagent Co., Ltd (Shanghai, China). Nitric acid (HNO₃), sodium hydroxide (NaOH), disodium, anhydrous ethanol (C₂H₅OH), sodium sulfate (Na₂SO₄), and *N*,*N*-dimethylformamide (DMF) were purchased from Shanghai Chemical Reagent Company (Shanghai, China). All chemicals were utilised without purification.

Synthesis of Bi₂O₂CO₃

The Bi₂O₂CO₃ was prepared by a simple hydrothermal method. First, 6 mmol of Bi(NO₃)₃·5H₂O were dissolved into 30 mL of nitric acid (1 M) in a 100 mL round-bottomed flask. 2 mmol citric acid was then dissolved into the solution and vigorously stirred for 10 min. 2 M NaOH aqueous solution was slowly dropped into the solution to adjust the pH value to 4. The mixture was heated at 180 °C for 24 h after having transferred to a 100 mL hydrothermal autoclave. The subsequent white precipitate was centrifuged and rinsed with deionised water/ absolute ethanol three times. The final Bi₂O₂CO₃ sample was obtained by drying the precipitate in a vacuum for 12 h at 60 °C (denoted as BO). Samples of BO were calcinated in a covered square crucible and heated at intervals of 3 °C min⁻¹ to 250-500 °C; the sample was then kept for 2 h in an atmosphere of hydrogen. The calcinated samples were denoted as BO-X (X =250, 300, 350, 400, 450, and 500 °C) consistent with the specific calcination temperature.

Synthesis of BO-X@PAN-Y

10 mg of PAN were added to 25 mL of DMF. A given mass (corresponding to $M_{PAN}: M_{BO-X} = 1:10, 1:100, 1:140, 1:180$) of BO-*X* was then dispersed into this solution with 4 h of stirring followed by ultrasonication for 1 h. The mixed solution was desiccated at 150 °C for 12 h to evaporate the *N*,*N*-dimethylformamide. The final nanocomposite was prepared by calcining the mixture at 350, 400, and 450 °C for 2 h in an N₂ atmosphere. The prepared samples were denoted as BO-*X*@PAN-*Y*(*Y* = 10, 100, 140, 180) consistent with the mass ratio of PAN to BO-*X*.

Characterisations

XRD (DX-2700, China) with Cu-K α radiation ($\lambda = 0.15406$ Å) was employed to investigate the crystalline phases in the 2θ region from 10° to 80° with the operating parameters of 40 kV and 150 mA. Images of the sample morphology were obtained via SEM (Zeiss Sigma 300, Germany) and TEM (FEI Tecnai G2 F20, Netherland). The chemical properties and structure of the catalyst surface were characterised by XPS (Thermo Scientific K-Alpha, USA) with Al K α X-ray ($h\nu$ – 1486.6 eV) radiation. The binding energy was calibrated with C 1s = 284.50 eV as the reference. TGA was carried out using a thermal gravimetric analyser (TGA5500, USA) at 10.0 °C min⁻¹ under a N₂ atmosphere (10 mL min⁻¹). UV-vis DRS were obtained using a UV-2600 (Shimadzu, Japan) and with BaSO₄ as a reference. The electrochemical workstation (CHI 660E, Shanghai Chenhua, China) was used to detect the TPR (I-t plots) with a conventional three-electrode configuration in a square quartz electrolytic cell under radiation by a 300 W Xe lamp at 20 s intervals. EIS was conducted with the conventional three-electrode system and recorded over a frequency range of 0.01 Hz-10⁵ Hz with an AC amplitude of 50 mV. Gauze platinum and silver chloride electrodes functioned as counter electrodes in the conventional three-electrode system. 0.5 M sodium sulfate solution was utilised as the electrolyte for the TPR. The mixed solution of potassium ferricyanide (0.1 M), potassium ferrous cyanide (0.1 M) and potassium chloride (0.1 M) was used as the electrolyte for electrochemical impedance experiments. Photoluminescence (PL) emission spectra were generated using the fluorescence spectrophotometer (FSL1000, Edinburgh Instruments Ltd, United Kingdom). The ESR signals of active species were collected on a spectrometer (Bruker EMXPLUS, Germany). Hydroxyl radicals ('OH) and positive holes (h⁺) were detected by dispersing the samples in a 50 mM solution of 5,5-dimethyl-1pyrroline N-oxide (DMPO) under visible-light illumination ($\lambda >$ 420 nm).

Photocatalytic activity test

The photocatalytic performance of the catalysts was estimated after removing TC or RhB from water. A 300 W Xenon lamp with a 420 nm pass filter was adopted as the light source. 50 mg of the prepared samples were dispersed in a quartz glass reactor containing 100 mL of aqueous TC (30 mg L⁻¹) or RhB (30 mg L^{-1}) water solution. The experimental temperature was controlled at room temperature by a circulating water bath system. To reach adsorption-desorption equilibrium, the mixture was continuously stirred in the dark for 30 min prior to photocatalysis. During irradiation, the reaction solution (5 mL) was sampled (5 mL of suspension) at fixed time intervals. The catalyst was removed by filtration through a hydrolases syringe filter (0.22 µm). The concentrations of TC and RhB were evaluated by UV-vis spectrophotometer (TU-1901, Persee, China). The characteristic absorption peaks of TC at RhB (357 and 554 nm, respectively) were used to quantify the extent of degradation. The removal rate (η) of the photocatalyst was calculated from eqn (1):

$$\eta (\%) = (1 - C/C_0) \times 100\%$$
⁽¹⁾

The concentrations of filtrates before and after irradiation are represented by the symbols C_0 and C, respectively.

The oxidation intermediate TC degradation products were measured by HPLC-MS (Agilent 1100 series, USA). The mobile phase employed a mixture of acetonitrile and formic acid (0.1%) with a flow rate of 0.135 mL min⁻¹ (injection volume and mass range: 5.0 μ L and 50–500 *m/z*, respectively; fragment and capillary voltage: 130 and 2800 V, respectively). Various trapping agents were added prior to the photocatalytic reaction to detect the existence of active free radicals.

Results and discussion

Structures and compositions of photocatalysts

Bi2O2CO3 and modified Bi2O2CO3 precursors were first synthesised using a facile hydrothermal method. The photocatalyst synthetic process is illustrated in Scheme 1. As shown in Fig. 1, the crystal structures of the Bi₂O₂CO₃, β-Bi₂O₃, pure α-Bi₂O₃, $Bi_2O_2CO_3/\beta$ - Bi_2O_3 , α - Bi_2O_3/β - Bi_2O_3 , and $Bi_2O_2CO_3/\alpha$ - Bi_2O_3 powders were analysed by XRD. Fig. 1a illustrates the XRD patterns of the Bi₂O₂CO₃ precursor prior to polymer modification and annealing at various temperatures. At calcination temperatures less than 250 °C, all the BO-250 diffraction peaks corresponded to the Bi2O2CO3 crystal planes (JCPDS no. 41-1488).27 Annealing at temperatures between 250 and 300 °C decoupled the C-O bonds of Bi₂O₂CO₃ corresponding to the gradual reduction of carbon elements and the conversion of $Bi_2O_2CO_3$ to β - Bi_2O_3 . When the calcination temperature was increased to 350 °C, a more prominent β-Bi₂O₃ peak appeared $(2\theta = 27.90^{\circ})$ (JCPDS no. 27-0050), suggesting a considerable increase in the β -Bi₂O₃ content in the composite. α -Bi₂O₃ manifested at 400 °C (JCPDS no. 71-2274), while at approximately 500 °C, the β-Bi₂O₃ crystal evolved into α-Bi₂O₃ following the rupture of double Bi₂O₃=Bi₂O₃ bonds.²⁸ The introduction of the polymer was also associated with comparable transitions in the crystal morphology with temperature (Fig. 1b). The observations of BO-400@PAN-140 indicated that the diffraction peak of $Bi_2O_2CO_3$ at $2\theta = 30.251^\circ$ reappeared after the polymer was added. Fig. 1c and d present the influence of the polymer to precursor ratio. The diffraction peak intensity of Bi2O2CO3 gradually enhanced with the upsurge of the polymer ratio, which may be caused by PAN coating on the outside of Bi₂O₂CO₃ to protect Bi₂O₂CO₃ from transformation during calcination,



Scheme 1 Schematic diagram of sample synthesis.

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Fig. 1 XRD patterns of (a) the $Bi_2O_2CO_3$ precursors and (b) polymermodified $Bi_2O_2CO_3$ precursors were calcined at various temperatures. (c and d) Polymer-modified $Bi_2O_2CO_3$ with various molar ratios (polymer/ $Bi_2O_2CO_3$) calcined at 400 °C.

while also indicating the formation of an $\alpha\text{-}Bi_2O_3/Bi_2O_2CO_3$ heterojunction (Fig. S1†).

Thermogravimetric analysis (TGA) analyses were performed to further quantify the phase transformation behaviour during the precursor calcination process.²⁹ Fig. S2† presents a relatively wide endothermal temperature range from 100 to 270 °C, with the highest peak emerging at 335 °C and a comparatively steady, stage-based performance between 270 and 320 °C. This phase transformation behaviour is consistent with the XRD results; for example, the Bi₂O₂CO₃ precursor is converted in a piecemeal manner to Bi₂O₂CO₃/ β -Bi₂O₃ and pure β -Bi₂O₃ at a specific intermediate temperature because of the thermal decomposition. β -Bi₂O₃ is then converted to β -Bi₂O₃/ α -Bi₂O₃ and α -Bi₂O₃ following calcination at higher temperatures.

XPS analyses were further conducted to investigate the surface chemical components and the elemental states. The survey spectra of BO-400 and BO-400@PAN-140 (Fig. 2a) indicate the presence of C, O, Bi, and N on the material surface. In Fig. S3,† the two peaks in the O 1s spectrum of BO-400@PAN-140 with binding energies at 530.70 and 529.11 eV correspond to metal carbonate and metal oxide, respectively. The N element is derived from the PAN molecules, while the C element of BO-400 originates from adventitious elemental carbon (Fig. S4†). The peak positioned at 288.90 eV is likely the result of carbon dioxide from the environment. In Fig. 2b, the two peaks in the Bi 4f spectrum of BO-400@PAN-140 with binding energies at 163.65 and 158.35 eV correspond to Bi $4f_{5/2}$ and Bi $4f_{7/2}$ of Bi³⁺, respectively.^{30,31}

In addition, the two peaks at 163.75 and 158.45 eV of Bi 4f are related to BO-400. In Fig. 2c, the C 1s spectra of BO-400@PAN-140 can be deconvoluted into three components at approximately 284.80, 286.60, and 288.90 eV. The peak consigned to 284.80 eV is associated with the C-C bonds of polyacrylonitrile, while the peak at approximately 286.60 eV results from the C-O of the adventitious carbon species. Finally, the peak near 288.90 eV corresponds to C=O, which is attributed to CO_3^{2-} in



Fig. 2 X-ray photoelectron spectroscopy (XPS) survey spectra: (a) high-resolution XPS spectra of Bi 4f; (b) BO-400 and BO-400@PAN-140 and C 1s (c); N 1s; (d) BO-400@PAN-140.

 $Bi_2O_2CO_3$. Fig. S5[†] presents the X-ray photoelectron spectroscopy (XPS) spectra of the carbon element following the introduction of the polymer and prior to calcination. The BOC@PAN-140 peak located at 286.30 eV persists and is unaffected by the association with BO-400@PAN-140. The BO-400@PAN-140C 1s binding energy at 288.90 eV demonstrates an obvious blue shift relative to BOC@PAN-140, which is likely attributed to the alteration of the chemical bond energy between the bismuth atom and surface carbon species of $Bi_2O_2CO_3$ the latter corresponding to the adventitious carbon. The emergence of the carbonate peak indicates that the polymer inhibits the transition of bismuth oxycarbonate to bismuth oxide in a hightemperature environment. As shown in Fig. 2d, the N1 s peaks at 398.40 and 399.79 eV can be ascribed to C–N and C==N in BO-400@PAN-140.³²

Fig. 3a-c presents the scanning electron microscopy (SEM) images of the surface morphologies of Bi2O2CO3, BO-400@PAN-140, and BO-400@PAN-10. Bi2O2CO3 can be seen to possess a layered structure with excellent structural homogeneity (Fig. 3a). As shown in Fig. S6a,[†] heating the Bi₂O₂CO₃ to 300 °C results in the aggregation of these nanosheets partially transforming them into nanometer-scale microspheres. Synthesizing this finding with the XRD data analysis, it is conjectured that these nanospheres are β -Bi₂O₃. Fig. S6b⁺ indicates that the lateral size of the nanosheet decreases following the introduction of PAN. A small number of tiny nanospheres are present between the nanosheet layers, deriving from the partial inhibition of the transformation from $Bi_2O_2CO_3$ to β -Bi₂O₃. As shown in Fig. 3b, the partial nanosheets were transformed into unique dendritic structures. The transformation from Bi2O2CO3 to α-Bi₂O₃ is the result of a solid reaction process.³³ The asproduced grain boundary presents a plausible reactive site for thermal decomposition. As a result of the polymer additive, the coated Bi₂O₂CO₃ transformed the a-Bi₂O₃/Bi₂O₂CO₃



Fig. 3 Scanning electron microscopy (SEM) images of (a) $Bi_2O_2CO_3$, (b) BO-400@PAN-140, and (c) BO-400@PAN-10. Transmission electron microscopy (TEM) images of (d) BO-400@PAN-140. (e) Elemental dispersive spectroscopic (EDS) mapping profiles of (f) Bi, (g) O, (h) C, and (i) N.

heterostructure at 400 °C. In addition, the mixing ratio of the polymer and Bi₂O₂CO₃ was found to significantly influence the final morphology of the photocatalyst. As shown in Fig. 3c, the extent of catalyst agglomeration increased with the polymer ratio. A mass ratio of Bi2O2CO3 to PAN of 1:10 corresponded to the formation of a dendritic microsphere surface scattered with small, irregular flakes. Reducing this mass ratio to 1:100 led to the formation of smaller dendritic structures with stacked nanosheets (Fig. S6c[†]). As shown in Fig. S6d,[†] a mass ratio of Bi₂O₂CO₃ to PAN of 180:1 resulted in a significant increase in the number of dendritic structures formed. Simultaneously, the small number of lamellar structures can be attributed to the inhibition of bismuth oxycarbonate transformation. A representative high-resolution transmission electron microscopy (HR-TEM) image of the BO-400@PAN-140 sample is shown in Fig. 3d, in which the shaded area indicated by the yellow arrow is Bi₂O₂CO₃. The degree of lattice spacing between Bi₂O₂CO₃ nanosheets is consistent with the (013) spacing of the Bi₂O₂CO₃ phase. The α -Bi₂O₃ phase is indicated by the pink arrow; the degree of lattice spacing in this region is consistent with crystal planes of α -Bi₂O₃ (200). These visualisations confirm that the BO-400@PAN-140 sample possesses a demonstrable heterostructure with a well-contacted interface.³⁴ As shown in Fig. 3e, the EDS (elemental dispersive spectroscopic) mapping of composite BO-400@PAN-140 was obtained for the same region of the sample. The sample was found to be composed of bismuth, oxygen, carbon, and nitrogen-further confirming the presence of the two phases which comprise the targeted heterostructure. The distributions of the elements presented in Fig. 3f-i confirm the presence and even distribution of all four elements from each of the two phases.

Photocatalytic activity

The photocatalytic degradation performance of the synthesised catalysts, with respect to both TC and RhB, was evaluated under visible-light radiation. Fig. 4 presents the photodegradation of TC (as followed by its change in concentration) as a function of time. As shown in Fig. 4a, the degradation rate of TC catalysed



Fig. 4 (a) Photocatalytic degradation curves of TC under visible-light irradiation ($\lambda > 420$ nm) with BO-400@PAN-Y (Y = 10, 100, 140, 180). (b) Photocatalytic degradation curves of RhB under visible-light irradiation ($\lambda > 420$ nm) with BO-400@PAN-Y (Y = 10, 100, 140, 180). (c) Photocatalytic degradation curves of TC with samples synthesised at various calcination temperatures. (d) The cycling runs in the photodegradation of TC over BO-400@PAN-140 composites.

solely by Bi₂O₂CO₃ and BO-400 was extremely slow compared to that which can be achieved in the presence of a polymer additive. Bi₂O₂CO₃ and BO-400 was associated with a TC removal efficiency of 40.9% and 49.84% within 60 min. As expected, the composites calcined at 400 °C exhibited an improved photocatalytic decomposition efficiency when compared with the conventional BO-400. Among these calcined samples, BO-400@PAN-140 possessed the highest photocatalytic performance (98.20%) for the degradation of TC within 60 min, which is 48.36% higher than that of BO-400. This increased photocatalytic activity may be attributed to the various crystal morphologies of Bi₂O₂CO₃ derived Bi₂O₃. The introduction of PAN inhibits the conversion of Bi₂O₂CO₃ to Bi₂O₃. The resulting $Bi_2O_2CO_3/\alpha$ - Bi_2O_3 heterojunction supports enhanced redox reaction. However, the degradation efficiency of TC decreased from 98.20 to 79.07% when the mass ratio of PAN was increased from 1:140 to 1:10 indicating that excessive PAN negatively impacts the photocatalytic degradation efficiency. Increased PAN may result in catalyst agglomeration, thereby limiting the contact area of the catalyst available to the pollutant. The TC removal efficiency of BO-400@PAN-10, BO-400@PAN-100, and BO-400@PAN-180 (within 60 min) were 79.07, 88.75, and 88.26%, respectively. The rate constant of the kinetic fitting curves (Fig. S7[†]) for Bi₂O₂CO₃, BO-400, BO-400@PAN-10, BO-400@PAN-100, BO-400@PAN-140, and BO-400@PAN-180 were 0.00633, 0.01009, 0.02371, 0.03455, 0.06358, and 0.03476 min^{-1} , respectively. The TC degradation rate constant over the α -Bi₂O₃/Bi₂O₂CO₃ heterojunction is nearly 6.3 times higher than over pure α -Bi₂O₃. Remarkably, the photocatalytic degradation activity of BO-400@PAN-140 was enhanced to levels comparable to the previously referenced best-in-class photocatalysts (Table 1). As shown in Fig. 4b, BO-400@PAN-140 can also enable the efficient photodegradation of RhB. The trend of RhB photodegradation efficiency with the PAN content of the

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Table 1 Photocatalytic degradation of TC by various photocatalysts

Photocatalysts	Light source	Dosage	Volume and concentration	Degradation efficiency	Ref.
CeO ₂ /C/Bi ₂ O ₂ CO ₃	Visible light	35 mg	100 mL; 20 mg L^{-1}	79.5% (90 min)	35
AgBr/5GO/Bi ₂ WO ₆	Visible light	40 mg	100 mL; 20 mg L^{-1}	84% (60 min)	34
$BiOCl_{0.9}I_{0.1}/\beta$ - Bi_2O_3	Simulated sunlight	20 mg	50 mL; 20 mg L^{-1}	82.4% (120 min)	36
OVs-BiOI/α-Bi ₂ O ₃	Visible light	50 mg	50 mL; 20 mg L^{-1}	84% (120 min)	37
β-Bi ₂ O ₃ /Bi ₂ O ₂ CO ₃	Simulated sunlight	100 mg	100 mL; 30 mg L^{-1}	93.79% (60 min)	21
BO-400@PAN-140 (α-Bi ₂ O ₃ /Bi ₂ O ₂ CO ₃)	Visible light	30 mg	100 mL; 30 mg L^{-1}	98.20% (60 min)	This work

photocatalyst is similar to that for TC. BO-400@PAN-140 possesses the highest photocatalytic performance (98.91%) for the degradation of RhB in 40 min. BO-400@PAN-10, BO-400@PAN-100, and BO-400@PAN-180 enabled the effective degradation of more than 90% of RhB within 40 min of visible-light illumination, compared to that by the original BO-400 (24.02%) and Bi₂O₂CO₃ (39.7%).

The rate of TC degradation was further investigated in relation to the calcination temperature of the photocatalyst (Fig. 4c). The catalyst calcination temperatures of 250, 350, and 400 °C corresponded to TC decomposition rates of 65.17, 83.10, and 49.84%, respectively, in the absence of the PAN additive. The decline in catalyst performance at elevated calcination temperatures may be understood in terms of the evolution of its crystal morphology *i.e.*, the complete conversion of Bi₂O₂CO₃ to β -Bi₂O₃ at 350 °C, while transforming completely into α -Bi₂O₃ 400 °C. The decomposition rates of TC achieved by BO-250@PAN-140, BO-350@PAN-140, and BO-400@PAN-140 were 72.19, 83.10, and 98.20%, based on a constant PAN mass ratio (1:140). The introduction of PAN inhibited the partial transformation of Bi2O2CO3 into α-Bi2O3 at elevated calcination temperatures, while the heterojunction formed by Bi₂O₂CO₃ and α-Bi₂O₃ improved the catalytic efficiency. As shown in Fig. S8,[†] the kinetic fitting curve rate constants for BO-250, BO-350, BO-400, BO-250@PAN-140, BO-350@PAN-140, and BO-400@PAN-140 were 0.01498, 0.02665, 0.01009, 0.01816, 0.02896, and 0.06358 min⁻¹, respectively. BO-400@PAN-140 exhibits the highest apparent rate constant, which is approximately 4.2, 2.4, 6.3, 3.5, and 2.2 times higher than those of BO-250, BO-350, BO-400, BO-250@PAN-140, BO-350@PAN-140, and BO-400@PAN-180, respectively. The experiments were repeated over five cycles under controlled visible-light irradiation to validate the reproducibility of the results.³⁸ The degradation rate of TC was maintained above 95% in each cycle (Fig. 4d). Out of all the possible permutations of PAN composition and calcination temperatures, BO-400@PAN-140 exhibited the most superior photocatalytic degradation performance in terms of both activity and stability. Fig. S9[†] clearly indicated that the prepared composites BO-400@PAN-140 shows a 45% mineralisation efficiency of TC within 100 min visible light irritation. This result confirmed that BO-400@PAN-140 could mineralise TC into intermediate compounds or CO₂ and H₂O.

Optical properties

As established in previous studies, both light absorption characteristics and energy band features are critical to the performances of the photocatalysts.³⁹ Fig. 5a presents the UV-vis diffuse reflectance spectra of the $Bi_2O_2CO_3$, BO-400, and BO-400@PAN-140 photocatalysts. $Bi_2O_2CO_3$ was found to absorb UV light limited by absorption edge at 390 nm. In contrast, BO-400, and BO-400@PAN-140 can absorb wavelengths in the visible-light region indicating their suitability for visible-light driven photocatalystic reactions. The absorption edges of the semiconductor photocatalysts BO-400 and BO-400@PAN-140 are 440 and 630 nm. The absorptive band edge of BO-400@PAN-140 is red-shifted relative to that of BO-400 indicating that the addition of the polymer inhibits the partial transformation of bismuth oxycarbonate.⁴⁰ Such heterojunction photocatalysts can enhance the visible light absorption while narrowing its bandgap, maximising the number of photogenerated carriers.

The bandgap structure strongly influences the photocatalytic activity; it can be evaluated by Tuac's equation:

$$\alpha hv = A(hv - E_g)^{n/2} \tag{2}$$

where α , h, v, A, and E_g are the absorption coefficient, Planck constant, light frequency, empirically determined constant, and the bandgap energy, respectively. The value of n, which depends on the transition characteristics of the semiconductor, is taken



Fig. 5 (a) UV-vis diffuse reflectance spectra of $Bi_2O_2CO_3$, BO-400, and BO-400@PAN-140 composites. (a) Tauc's plots of $Bi_2O_2CO_3$ and BO-400. (b) Transient photocurrent response (TPR) curves of BO-400@PAN-Y. (c) Electrochemical impedance spectroscopy (EIS) spectra of BO-400 and BO-400@PAN-140. (d) PL spectra of BO-400 and BO-400@PAN-Y composites.

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as 4 in this context based on the indirect transformation of $Bi_2O_2CO_3$ to α - Bi_2O_3 .⁴¹ The band gaps (Fig. 5a1) of α - Bi_2O_3 and $Bi_2O_2CO_3$ were determined to be 2.96 and 2.65 eV.

The band edge positions of $\rm Bi_2O_2CO_3$ and $\alpha{\rm -Bi_2O_3}$ can be predicted theoretically based on electronegativity. The valence band (VB) and conduction band (CB) potentials of the asprepared, charge-neutral photocatalysts were determined as follows. 41,42

$$E_{\rm VB} = X - E_{\rm e} + 0.5E_{\rm g} \tag{3}$$

$$E_{\rm CB} = E_{\rm VB} - E_{\rm g} \tag{4}$$

where $E_{\rm g}$ is the bandgap energy of the semiconductor; $E_{\rm VB}$ corresponds to the valence band edge potential; $E_{\rm CB}$ corresponds to the conduction band edge potential; *X* is the absolute electronegativity of the semiconductor, which is the geometric mean of the absolute electronegativity of the constituent atoms (*X* for Bi₂O₂CO₃ and α -Bi₂O₃ is 6.36 and 6.23 eV, respectively);⁴³ and $E_{\rm e}$ is the energy of free electrons (~4.5 eV).

Based on the above formula and data, the VB and CB edge potentials of $Bi_2O_2CO_3$ are estimated to be 3.24 and 0.28 eV, respectively, while the VB and CB edge potentials of α -Bi₂O₃ are estimated to be 3.05 and 0.40 eV. To further validate this inferred band structure, the valence band edges of $Bi_2O_2CO_3$ and α -Bi₂O₃ were characterised based on the valence-band XPS (VB-XPS) measurements. As shown in Fig. S10,[†] the valence band edges of original $Bi_2O_2CO_3$ and α -Bi₂O₃ are 3.22 eV and 3.03 eV, which are comparable to the results of the theoretical calculation. These energy band structures are also presented in Table S1.[†]

The charge transfer and separation efficiency of photogenerated electron-hole pairs are crucial to photocatalytic performance, as reflected by the transient photocurrent response (TPR).44 The TPR profiles shown in Fig. 5b demonstrate that BO-400 and BO-400@PAN-Y were stable in response to each irradiation, while also confirming that the photocurrent response of BO-400@PAN-Y is higher than that of BO-400, which may be derived from the formation of the $Bi_2O_2CO_3/\alpha$ - Bi_2O_3 p-n heterojunction. The existence of this heterojunction can also improve both the transferability and separability of the photogenerated carriers. BO-400@PAN-140 exhibited the highest TPR; irradiation by a medical cold light source generated a rapid photocurrent that reached a steady-state over a short interval. Electrochemical impedance spectroscopy (EIS) was employed to further validate interfacial charge transfer behaviour.45 As illustrated in Fig. 5c, the arch radius of the heterojunction catalyst BO-400@PAN-140 impedance curve is smaller than that of BO-400, BO-400@PAN-10, BO-400@PAN-100, and BO-400@PAN-180, further demonstrating that the interaction between α-Bi₂O₃ and Bi₂O₂CO₃ can effectively reduce the charge transfer resistance at the phase interface. These findings confirm that BO-400@PAN-140 possesses significantly improved interface charge transfer and electron-hole pair separation efficiencies. The separation and recombination of photogenerated carriers can be further detected from the photoluminescence (PL) spectra.⁴⁶ The PL spectra of BO-400 and

BO-400@PAN-140 photocatalysts are presented in Fig. 5d, based on an excitation wavelength at 375 nm. BO-400 exhibits a strong emission at 530 nm due to carrier recombination under optical radiation. Relative to BO-400, the BO-400@PAN-*Y* photocatalysts exhibited a noticeably quenched PL signal, indicating that the formation of α -Bi₂O₃ and Bi₂O₂CO₃ heterojunction inhibits the direct recombination of photogenerated electrons and holes. The PL intensity of the BO-400@PAN-*Y* nanocomposites decreases as the ratio of PAN is reduced, corresponding to enhanced electron mobility in the conjugated system. In summary, the photogenerated electron/hole between Bi₂O₂CO₃ and α -Bi₂O₃ favors electron transfer and, by extension, the photocatalytic degradation rate.

Possible degradation mechanism

To better characterise the photocatalytic mechanism of BO-400@PAN-140, a series of free radical trapping experiments were conducted to systematically identify the active radicals involved in the decomposition of organic pollutants. L-Ascorbic acid (L-AA), ethylene diamine tetraacetic acid (EDTA), and *tert*butyl alcohol (TBA) were employed as the scavenging reagents to quench peroxide radicals (' O_2^-), holes (h⁺), and hydroxyl radicals ('OH), respectively.⁴⁷ As shown in Fig. S11,† TC degradation was almost unaffected after the addition of 1 mM L-AA. In contrast, the addition of TBA and EDTA significantly affected the degradation of TC, indicating that 'OH and h⁺ played a major role in the removal of TC over α -Bi₂O₃/Bi₂O₂CO₃ heterojunction catalysts.

EPR (Electron Paramagnetic Resonance) experiments were conducted to further validate the active species involved in the photodegradation of TC over BO-400@PAN-140. The capture agents for h⁺ and 'OH were TEMPO and DMPO, respectively.¹⁹ As indicated in Fig. 6a, the signal intensity of h⁺ following 5 and 10 min of light exposure is weaker than without any exposure, implying that the photocatalyst is conducive to the separation and migration of photogenerated electrons/holes under irradiation. Fig. 6b indicates the absence of the 'OH signal under dark conditions. However, the signal intensity of 'OH was shown to gradually increase with the increasing light irradiation.⁴⁸ BO-400@PAN-140 significantly enhances molecular oxygen activation to generate 'OH, which is derived from strong interactions engendered in the Bi–N bonds at the interface of the photocatalyst and PAN additive.^{49,50} These EPR results



Fig. 6 Electron paramagnetic resonance (EPR) spectra of the (a) DMPO- h^+ , and (b) DMPO--OH adducts were recorded with the BO-400@PAN-140 under the dark and light irradiation.

further validate the generation of h^+ and 'OH radicals under visible light, which is also consistent with the results of radical trapping experiments.

Possible tetracycline degradation pathway

HPLC-MS results highlight the key intermediate products and possible reaction pathways for photocatalytic degradation for TC over BO-400@PAN-140 (the corresponding molecular formulas and structure of which are presented in Fig. S12[†]). Using insights from the relevant literature and detected intermediates, Fig. 7 illustrates two possible TC degradation pathways.⁵¹ The active species, which consist of 'OH, and h⁺, can effectively attack partially unstable TC groups including amine, double-bond, and phenolic groups, resulting in the cleavage of such functional groups as the photocatalytic reaction progresses.52 The base peak chromatogram of untreated solution exhibits its main peak at m/z = 445, corresponding to the tetracycline molecule. Pathway I is characterised by a methyl of the amino group on T1 gradually colliding into the minor molecule T2 (m/z = 431) by h⁺ attack of the methyl and aldehyde species. Subsequent 'OH degradation of the methyl species on the tertiary amine leads to the formation of T5 (m/z = 417).⁵³ The transformation from T5 to T6 (m/z = 345) is driven by the cleavage of the acylamido group and open-ring reactions.54 Pathway II is characterised by the attack of T1 (m/z = 445) by 'OH (formed from H₂O) to produce the product T3 (m/z = 428).⁵⁵ Given its high electron density, the cleavage of the C-N bond can be attributed to the formation of the intermediate product



Fig. 7 Intermediate products and proposed degradation pathway of TC.

T4 (m/z = 399). T7 and T8 were transformed into small organic molecular byproducts (m/z = 118, m/z = 164, and m/z = 136) following a series of open-ring reactions and the forfeiture of specific groups. These small organics were ultimately converted to CO₂ and H₂O by an attack of 'OH and h⁺.

The relative band edge positions of the two materials determine the direction of migration of photogenerated electrons and holes on the p-n heterojunction photocatalyst. The likely mechanism for the enhancement of photocatalytic degradation activity follows from the preceding analysis and energy band calculations.^{56,57} In contrast with the CB and VB positions of pure Bi₂O₂CO₃ and α-Bi₂O₃, the p-n heterojunction photocatalyst exhibits nested band energy levels (left of Scheme 2), which is unfavourable for the parting and transmission of photogenerated carriers. The Fermi energy level (E_f) of α -Bi₂O₃ is close to its valence band, while Bi₂O₂CO₃ approximates its conduction band.^{58,59} The CB and VB locations of α -Bi₂O₃ and Bi₂O₂CO₃ are shifted as a result of the formation of the p-n heterojunction structure. The E_f of p-type α -Bi₂O₃ would increase, while that of n-type Bi₂O₂CO₃ decrease, ultimately reaching a new equilibrium to form a new staggered band structure.

Following this transformation of the energy band structure, the conduction band position of $Bi_2O_2CO_3$ is lower than that of α -Bi₂O₃. This optimised energy band structure favours the departure and migration of photocatalytic carriers. The inner electron field forms at the interface of n-type $Bi_2O_2CO_3$ and ptype α -Bi₂O₃. Consistent with the staggered band structure of the composite photocatalyst, a promising photocatalytic degradation mechanism is proposed—as illustrated to the right of Scheme 2. Under visible-light irradiation, photogenerated carriers emerge from the α -Bi₂O₃ and Bi₂O₂CO₃ phases. The optimised energy band structure is conducive to the rapid transfer of photogenerated electrons from the α -Bi₂O₃ conduction band to the Bi₂O₂CO₃ conduction band.

As electrons migrate between the conduction bands of two components, the holes in the valence band of Bi₂O₂CO₃ will rapidly transfer to that of α -Bi₂O₃, ultimately forming a p-n heterojunction between α-Bi2O3 and Bi2O2CO3 to effectively separate electron-hole pairs. Simultaneously, the heterojunction-induced electric field inhibits the recombination of the electrons and holes at the interface of the two semiconductors. The charges migrate to the surface of the catalysts with the dissolved oxygen and OH⁻ in water, enabling the continuous formation of essential active groups. The conduction band potentials of Bi2O2CO3 and α-Bi2O3 are both higher than the redox potential of $O_2/O_2^{-}(-0.33 \text{ eV})$.⁶⁰ Therefore, dissolved oxygen in water cannot be reduced to O_2^{-} , having gained electrons in the conduction band.

The redox potential of O_2/H_2O_2 is 0.695 eV,^{61,62} which is higher than the CB potential of Bi₂O₂CO₃. Consequently, O₂ can initially be reduced to H₂O₂, further enabling the capture of electrons to form 'OH. The α -Bi₂O₃ possesses a higher valence band potential than the redox potential of OH⁻/'OH (2.40 eV).⁶³ Holes in the valence band of α -Bi₂O₃ oxidise OH⁻ to 'OH, while also migrating to the surface of the catalyst to promote the enhanced oxidation of TC. Together, 'OH and h⁺ function as



Scheme 2 A possible mechanism for pollutant degradation and electron flow with the BO-400@PAN-140 photocatalyst

strong oxidizing species for the degradation of TC contaminant molecules on the surface of the photocatalyst and the superior degradation performance (under visible-light irradiation) of the p–n heterojunction photocatalyst relative to that of a single Bi₂O₂CO₃ and α -Bi₂O₃ originates from its numerous mechanistic advantages.

Conclusions

A α-Bi₂O₃/Bi₂O₂CO₃ p-n heterojunction photocatalyst has been successfully synthesised via a facile annealing method. The a-Bi₂O₃/Bi₂O₂CO₃ p-n heterojunction demonstrated excellent photocatalytic performance for the degradation both of antibiotic (TC) and dye (RhB) contaminants, which can be attributed to the existence of a suitable energy band structure enabling the rapid separation of photogenerated carriers. Following only 60 min of visible-light irradiation, TC degradation efficiency of 98.2% was achieved using the BO-400@PAN-140 photocatalyst. This degradation efficiency was maintained above 95% after the completion of five cycles. Free radical scavenging tests and electron paramagnetic resonance (EPR) results demonstrated h⁺ and 'OH radicals as the primary active species in photocatalytic degradation. The HPLC-MS results also indicated that the TC molecules can be effectively decomposed through two pathways. For this reason, our proposed methodology for the synthesis of a-Bi2O3/Bi2O2CO3-based p-n heterojunctions with high contaminant photodegradation efficiency can make a significant impact on improving the readiness of photocatalytic water purification technologies.

Author contributions

Baikang Zhu: writing—review & editing, project administration, funding acquisition. Debin Song: conceptualization, methodology, visualization, writing—original draft. Lihui Chen: writing—review & editing. Qingguo Chen: writing—review & editing. Chunyang Zhai: writing—review & editing. Bohong Wang: writing—review & editing. Jiří Jaromír Klemešf: writing review & editing, funding acquisition. Hengcong Tao: conceptualization, methodology, writing—review & editing, supervision, funding acquisition. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

The authors declare no competing financial interest.

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