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Efficient photo-oxidation of bisphenol a and tetracycline through sulfur-doped g-C₃N₄/CD heterojunctions†

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The synergistic effects of sulfur-doped g-C₃N₄ (SCN) and carbon dots (CD) in nanocomposite photocatalysts were explored for the degradation of contaminants of emerging concern (CECs), particularly phenolic pollutants such as bisphenol A (BPA) and tetracycline (TC). Various SCN/CD nanocomposites were synthesized via thermal polymerization by mixing with different CD concentrations with thiourea (1, 2, 3, and 4 by wt%), denoted as SCN/CD1, SCN/CD2, SCN/CD3, and SCN/CD4, respectively. The embedded CD functions as an intrinsic surface modifier on the surface of the SCN which facilitated the suppression of electron–hole recombination and promoted photocatalytic activity. Among the synthesized catalysts, SCN/CD3 exhibited remarkable efficiency, degrading a 50-ppm solution of BPA to 92.5% and TC to 90.7% within 60 minutes, utilizing 50 mg of catalyst. Moreover, SCN/CD3 demonstrated exceptional reusability over five cycles without significant degradation in efficiency. Radical scavenging experiments identified holes (h⁺) and superoxide radicals (O₂^{•−}) as the primary radical species responsible for pollutant degradation. This work highlights the potential of SCN/CD composite photocatalysts in solving water pollution concerns by elucidating a promising photocatalytic degradation process for CECs.

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1. Introduction

The escalating global demand for water resources has accentuated the need for effective water and wastewater treatment technologies. However, despite progress in traditional treatment methods, the prevalence of contaminants of emerging concern (CECs) poses a formidable challenge.^{1,2} These CECs, encompassing a spectrum of pollutants from pharmaceuticals³ to endocrine disruptors, persist in water sources, posing significant threats to both the environment and human health.

Among the myriad of CECs, bisphenol A (BPA)^{4,5} and tetracycline (TC)^{6–8} stand out due to their widespread presence and potential adverse effects. BPA, a ubiquitous component of plastic products,⁹ and TC, a common antibiotic, are prevalent in various industrial effluents and municipal wastewater streams. Alarming, the concentrations of TC in pharmaceutical wastewater can soar to staggering levels, reaching up to 1077 mg L^{−1},^{10,11} while BPA concentrations in landfill leachate and industrial effluents range from 0.37 to 17.2 mg L^{−1}.^{12,13} Such elevated concentrations underscore the urgency to devise efficient remediation strategies to mitigate their detrimental impacts.

Photocatalysis stands as a beacon of hope in the realm of environmental remediation and clean energy production, offering a green and sustainable solution to pressing global challenges.¹⁴ Graphitic carbon nitride (g-C₃N₄)¹⁵ is one of the several types of photocatalysts that has drawn interest since it is devoid of metals and reacts to visible light making it appropriate for a variety of uses, such the evolution of hydrogen or oxygen,^{16,17} CO₂ reduction,¹⁸ and environmental remediation.¹⁹ However, bulk g-C₃N₄ faces limitations including high defect density, fast charge recombination, and low specific surface area.²⁰ To address these challenges, researchers

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have explored sensitized doping as a strategy to boost the photocatalytic activity of $g\text{-C}_3\text{N}_4$.²¹ Structural defects are created in the $g\text{-C}_3\text{N}_4$ lattice by adding non-metallic elements like sulphur, which optimises the electron distribution and changes the band gap location.²² This facilitates charge transfer at the material interface, expands light absorption range, and promotes efficient electron-hole separation, thus enhancing overall efficiency. Sulfur-doped $g\text{-C}_3\text{N}_4$ (SCN) exhibits superior photocatalytic activity compared to other dopants, owing to its improved specific surface area, enhanced light utilization rate, and superior efficiency of electron-hole separation.^{23,24} Additionally, it serves as an excellent carrier in heterojunction structure construction, facilitating charge separation and providing additional binding sites for pollutant adsorption.

Furthermore, the presence of carbon dots (CD) in $g\text{-C}_3\text{N}_4$ nanocomposites can facilitate interfacial charge transfer and separation through heterojunction formation, leading to enhanced photocatalytic efficiency.^{25–30} The intimate contact between CD and $g\text{-C}_3\text{N}_4$ promotes efficient electron transfer across the interface, facilitating rapid redox reactions and improving overall catalytic activity.³¹ Additionally, the presence of CD can alter the band structure of $g\text{-C}_3\text{N}_4$, leading to a red shift in the light absorption spectrum and enabling utilization of a broader range of solar radiation for photocatalysis.³²

This study delves into enhancing the photocatalytic efficacy of $g\text{-C}_3\text{N}_4$ by sulfur doping and incorporating CD into its structure. Through systematic investigation, we aim to elucidate the mechanisms underlying the superior performance of $g\text{-C}_3\text{N}_4$ as a photocatalyst, particularly focusing on sulfur doping effects. Our efforts contribute to advancing the understanding of these materials for environmental remediation purposes. Moreover, we explore the potential of SCN/CD nanocomposites in degrading CECs such as BPA and TC, crucial for sustainable environmental cleanup initiatives. This research underscores the significance of tailored modifications in $g\text{-C}_3\text{N}_4$ to optimize its photocatalytic activity, paving the way for efficient degradation of CECs and promoting environmentally friendly remediation strategies.

2. Experimental section

2.1. Materials and method

The chemical reagents, namely thiourea ($\text{CH}_4\text{N}_2\text{S}$: $\geq 99.0\%$), urea ($\text{CO}(\text{NH}_2)_2$: 99.0%), citric acid ($\text{C}_6\text{H}_8\text{O}_7$: $\geq 99.5\%$), bisphenol A ($\text{C}_{15}\text{H}_{16}\text{O}_2$: 99.0%), tetracycline ($\text{C}_{22}\text{H}_{24}\text{N}_2\text{O}_8$: $\geq 95.0\%$), and ethanol ($\text{C}_2\text{H}_6\text{O}$: 98.0%) were purchased from Sigma-Aldrich, with excellent analytical quality. They were used just as received, with no further refinement. Every solution was prepared using deionized water as the medium.

2.2. Synthesis of CD

A hydrothermal process was employed to create CD. Initially, ultrasonication was used for two hours to dissolve 1.0 g of citric acid and 2.0 g of urea in deionized water. After that, the mixture was placed inside a 150 mL Teflon-lined stainless-steel autoclave and heated to 200°C for 8 hours to undergo hydrothermal treatment. To get rid of big particles, the black solution was centrifuged for 30 minutes after it cooled to normal temperature and ultimately, the supernatant was gathered.³³

2.3. Synthesis of SCN/CD

The CD/SCN photocatalyst was synthesized using a thermopolymerization method. In the standard procedure, 2.0 g of thiourea was mixed with a specific volume of CD solution in 10 mL of deionized water. Following the evaporation of the solution at 70°C , the resulting powder underwent heating to 550°C for 3 hours, with a gradual increase in temperature at a rate of 2.8°C per minute. The resultant CD/SCN powder was obtained through meticulous grinding and filtration. Different SCN/CD samples were prepared with varying CD contents (1%, 2%, 3%, and 4% by weight), designated as SCN/CD1, SCN/CD2, SCN/CD3, and SCN/CD4, respectively. Sulphur atoms entered the $g\text{-C}_3\text{N}_4$ network *in situ* during the thermal polycondensation process when SCN samples were synthesised without the addition of CD solution.³⁴ The schematic illustration of the synthesis procedure is shown in Fig. 1.

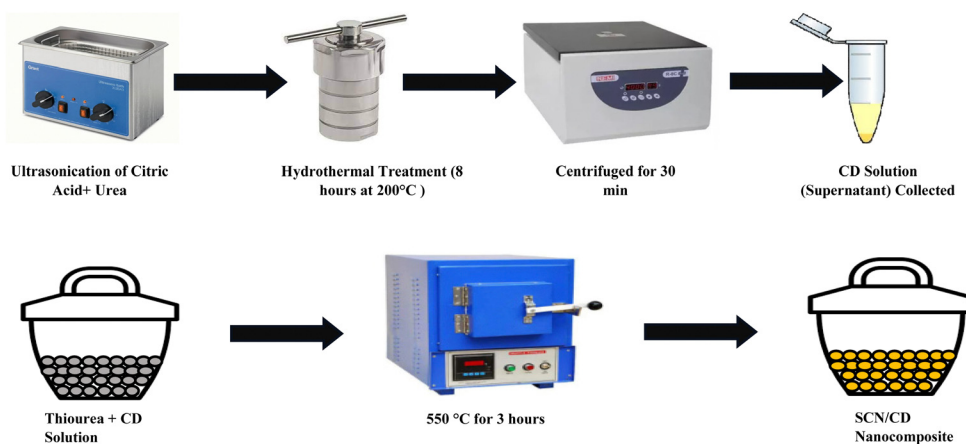


Fig. 1 Schematic representation of synthesis process of CD and SCN/CD.



3. Characterizations

3.1. X-ray diffraction (XRD)

XRD analysis was conducted within the angular range of 5° to 60° , offering valuable information on the structural properties of the CD, SCN and their combined nanocomposite under investigation. In Fig. 2, CD exhibited a broad hump centred at 21.8° on the (002) plane, indicating the presence of an amorphous carbon phase. Pure SCN displayed a minor peak at 12.8° implying the presence of regularly recurring units of the heptazine structure on the (100) plane. The significant peak at 27.5° indicated the layered arrangement of π -conjugated aromatic structures of g- C_3N_4 layers on the (002) plane.³⁵ Analysis of the SCN/CD nanocomposite revealed peaks resembling those of pure SCN, with no discernible CD peaks. A slight shift towards lower angles and a decrease in peak height were observed, attributed to CD amorphous nature, indicating some alteration due to lattice distance augmentation. Notably, XRD patterns remained consistent across varying CD concentrations, suggesting minimal influence on the lattice structure of SCN. This observation underscores the stability of SCN's lattice structure despite changes in CD concentration, as evidenced by uniform XRD profiles across the investigated CD concentration range.³⁶

3.2. Fourier transform infrared spectroscopy (FTIR)

The surface functionality of prepared nanocomposites was analysed using FTIR as shown in Fig. 3. In SCN spectra a broad peak in the region of $3050\text{--}3400\text{ cm}^{-1}$ was observed and identified as the N-H stretching vibration in terminal amino groups ($-\text{NH}_2$ or $=\text{NH}$ groups). The peaks at 1636 cm^{-1} and 1538 cm^{-1} indicated the C=N stretching vibration of the triazine unit, while peaks at 1400 cm^{-1} , 1315 cm^{-1} , and 1206 cm^{-1} corresponded to the C-N stretching. Additionally, the out-of-plane stretching of the tri-s-triazine unit was identified at 815 cm^{-1} .³⁷ Vibrations associated with the S-related group were minimal in doped g- C_3N_4 , possibly due to the low sulfur content and overlapping with C-N vibration peaks. In the

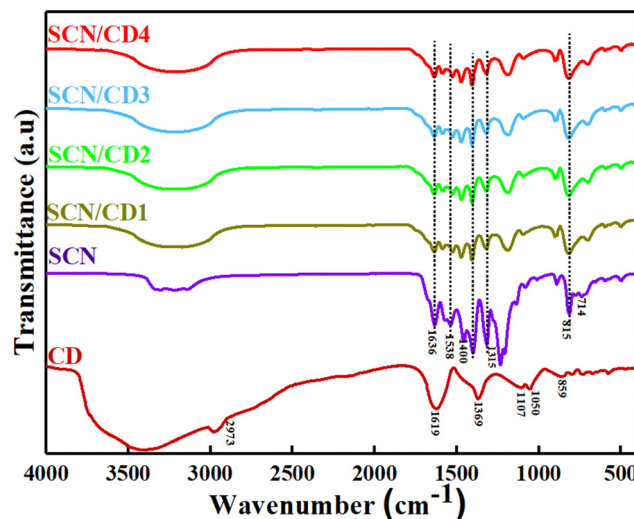


Fig. 3 FTIR patterns of CD, SCN and SCN/CD nanocomposites.

FTIR spectrum of CD, the broad hump at 3407 cm^{-1} were attributed to O-H stretching, with peaks at 2973 cm^{-1} and 1619 cm^{-1} associated with C-H stretching and C=C stretching, respectively. Peaks at 1369 cm^{-1} and 1050 cm^{-1} corresponded to C-N stretching vibrations.³⁸ Notably, in the SCN/CD spectrum, significant changes in the characteristic bands of SCN were observed, indicating potential interfacial interactions between CD and SCN. The FTIR spectra for a series of nanocomposites with varying concentrations of CD (1%, 2%, 3%, 4%) exhibited no discernible changes, indicating that the spectra remained consistent across different CD concentrations. This implies that the FTIR characteristics of the nanocomposites were not substantially changed by the addition of CD.

3.3. X-ray photoelectron spectroscopy (XPS)

XPS analysis of both the SCN and SCN/CD samples confirmed the presence of carbon (C), nitrogen (N), oxygen (O), and sulfur (S) components as shown in Fig. 4. Notably, the relative ratio of oxygen was higher in the SCN/CD3 sample, possibly due to absorbed water and oxygen by the CD. The C 1s transition displayed peaks at 287.2 eV , indicating sp^2 N-C=N bonds, and at 284.3 eV , associated with aromatic C atoms in the s-triazine ring. In the N 1s region, peaks were detected at 397.6 , 398.5 , 400.3 , and 403.3 eV , representing various nitrogen-containing groups such as sp^2 hybridization of localized nitrogen, tertiary N-(C)₃ groups, amino groups with hydrogen, and pi excitations. Peaks observed in the O 1s region at 530.6 eV and 532.6 eV likely stemmed from O-C/O-H and O-C components, potentially due to absorbed water and oxygen. Additionally, a distinct peak at 164 eV indicated the presence of a C-S bond, suggesting sulfur incorporation into the lattice structure, possibly through substitution with lattice nitrogen.^{32,39}

3.4. Ultraviolet-visible diffused reflectance spectroscopy (UV-DRS)

The optical absorption characteristics of the samples were probed using UV-visible diffuse reflectance spectroscopy. The

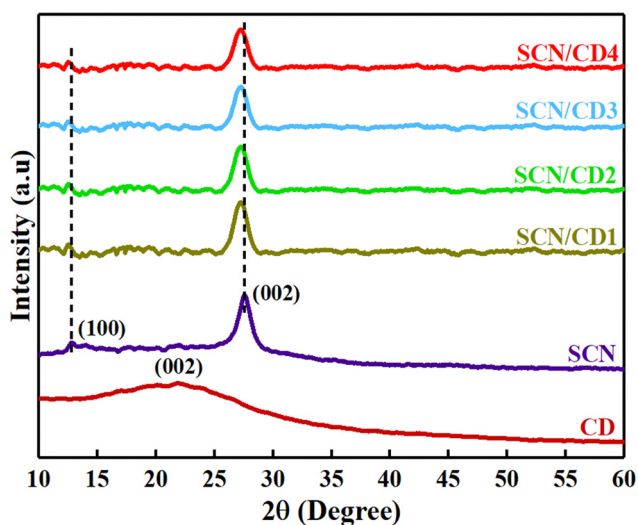


Fig. 2 XRD patterns of CD, SCN and SCN/CD nanocomposites.



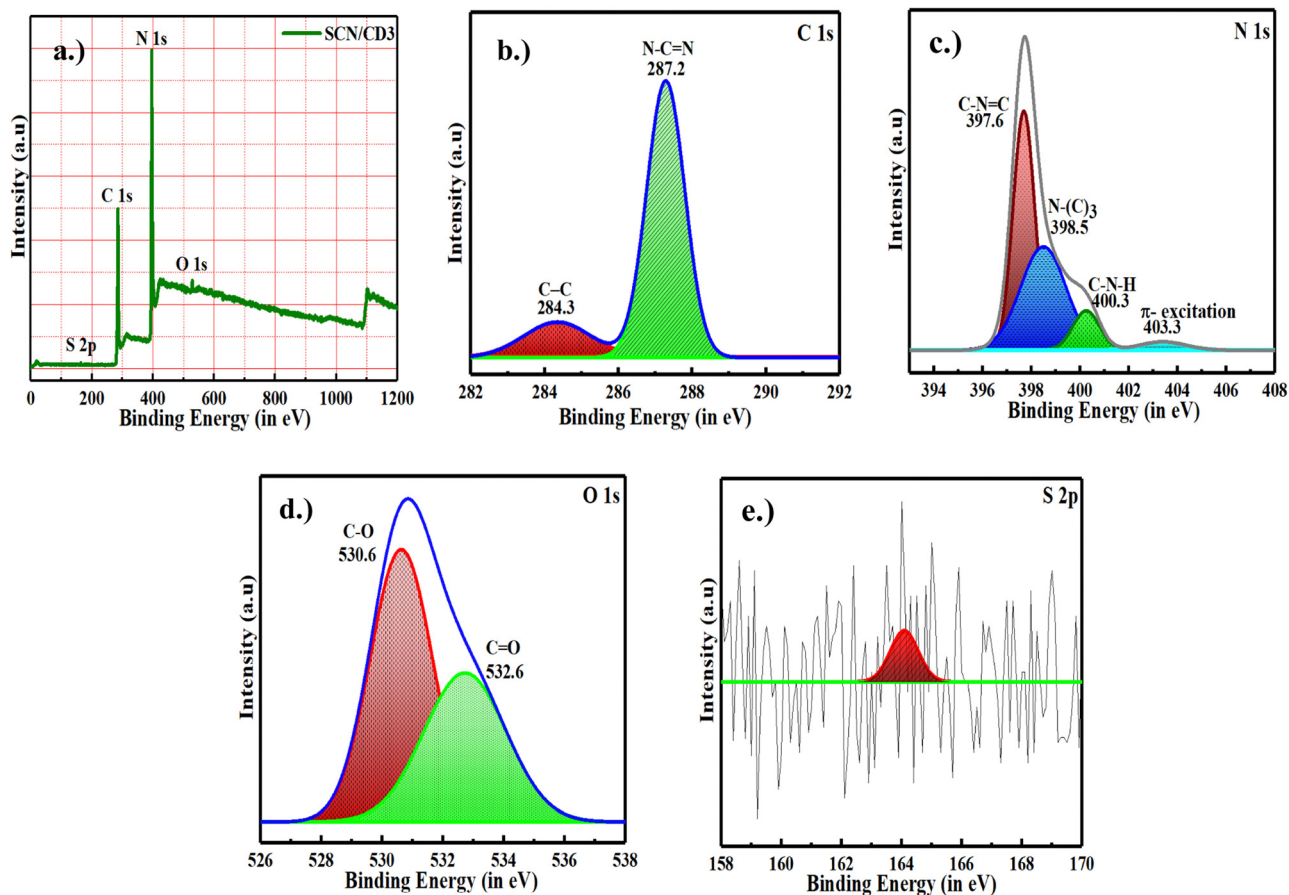


Fig. 4 (a) XPS spectra of SCN/CD3 nanocomposite. High resolution spectra of (b) carbon, (c) nitrogen (d) oxygen and (e) sulfur.

absorption onset for pristine was found at approximately 470 nm, while for both sulfur-doped carbon nitride (SCN) and SCN/CD nanocomposites, it shifted to around 500 nm as shown in Fig. 5(a). Interestingly, the incorporation of cyclodextrin (CD) into SCN led to a redshift in the absorption edge of the SCN/CD

nanocomposite, along with an enhancement in light absorption within the visible spectrum. The semiconductor bandgap was determined using the Tauc plot method, where the absorption coefficient (α) was plotted against the photon energy ($h\nu$). Fig. 5(b) analysis revealed a decrease in the bandgap from

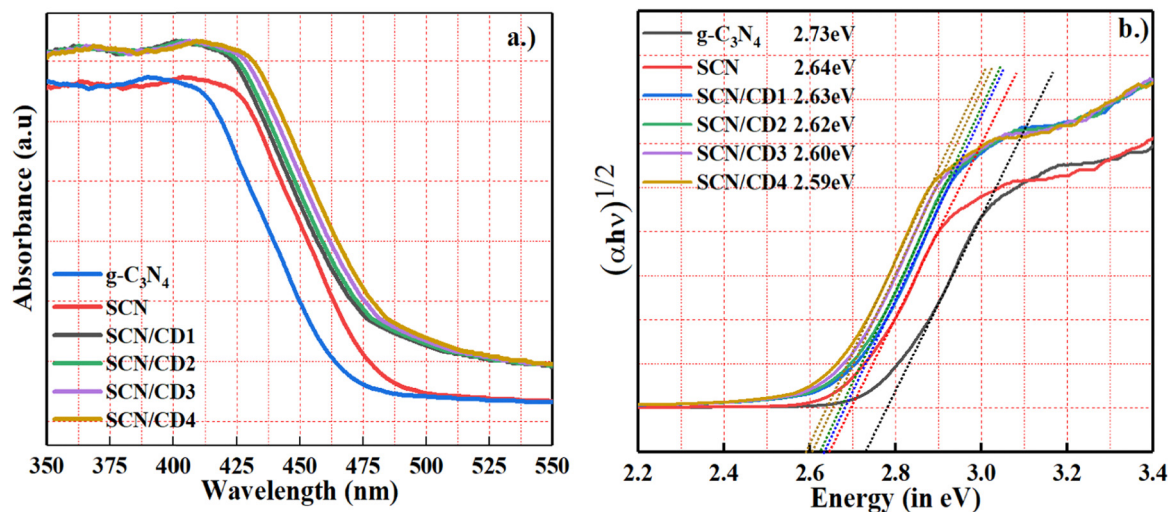


Fig. 5 (a) UV-vis diffuse reflectance of g-C₃N₄, SCN, SCN/CD1, SCN/CD2, SCN/CD3 and SCN/CD4 (b) Tauc plot representing the bandgap energies of g-C₃N₄, SCN, SCN/CD1, SCN/CD2, SCN/CD3 and SCN/CD4.



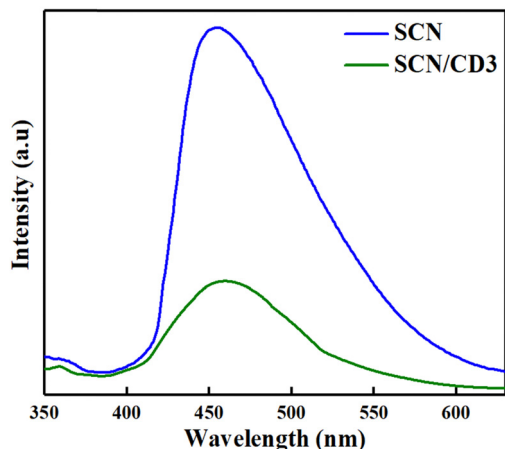


Fig. 6 PL spectra of synthesised nanocomposite SCN and SCN/CD3.

2.73 eV for g-C₃N₄ to 2.64 eV for SCN, indicating that SCN has a superior ability to harness visible light compared to pristine g-C₃N₄. Moreover, with the introduction of CD into SCN, the bandgap decreased further with increasing CD concentration, suggesting that CD incorporation marginally affects the nanocomposite's bandgap, potentially indicating CD's primary localization on the SCN surface.⁴⁰ Despite this, the narrowed bandgap shows promise for improving visible light absorption, facilitating the generation of photoinduced carriers.

3.5. Photoluminescence (PL) spectroscopy

PL spectroscopy was employed to examine the dynamics of electron-hole pair separation and recombination in SCN and SCN/CD3. With an excitation wavelength of 325 nm, SCN exhibited a distinct emission peak at approximately 455 nm, whereas SCN/CD3 displayed a slight red shift, manifesting a peak around 460 nm as shown in Fig. 6. Notably, the emission intensity of SCN/CD3 was notably lower than that of pristine SCN, indicating a significant suppression of electron-hole pair recombination within the composite photocatalysts.⁴¹ This suppression suggests efficient charge separation, facilitated by the combination of CDs with SCN, which accelerates electron transfer while hindering the recombination process, ultimately enhancing photocatalytic performance.

3.6. Electrochemical impedance spectroscopy (EIS)

The Nyquist plots generated from EIS data were utilized to assess the interfacial charge transfer characteristics of the synthesized photocatalysts. Typically, the arc radius observed in Nyquist plots corresponds to the reaction rate occurring at the working electrode. A smaller arc radius indicates a lower resistance electrode, suggesting efficient separation of electron-hole pairs and an adequate supply of electrons for the photodegradation reaction. In Fig. 7, it is observed that the SCN exhibited a larger arc radius compared to SCN/CD3. The observation implies that SCN/CD3 possesses the lowest resistance for charge transfer, indicating more effective separation of photoinduced charges.^{42,43} Such enhanced charge

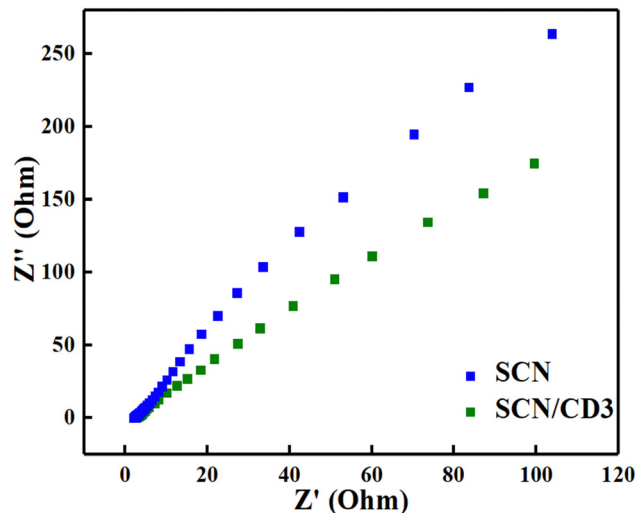


Fig. 7 EIS Nyquist plots of SCN and SCN/CD3.

separation is beneficial for improving photocatalytic performance, as it facilitates efficient utilization of photogenerated charge carriers.

3.7. Scanning electron microscope (SEM)

The morphology, structure, and elemental distribution of a nanocomposite were investigated using SEM. Fig. 8 depicts SEM images of both pure SCN and SCN/CD nanocomposite. The surface of SCN comprises flat and stratified nanosheets arranged in layers. In contrast, in the SCN/CD nanocomposite, the SEM images reveal the presence of small spherical CD particles dispersed on the surface of the SCN layered structure. Importantly, these CD particles are not merely adhering to the surface but are embedded within the SCN surface itself, effectively acting as intrinsic surface modifiers. This embedded configuration ensures intimate contact between the CD particles and the SCN material, enhancing their catalytic performance and efficiency in the photocatalytic degradation process. Also, the sheet-like morphology providing large surface area further facilitates the adsorption of organic molecules, thereby increasing the proportion of catalytically active sites on the surface. This enhancement contributes to the photocatalytic degradation process.

3.8. Energy dispersive X-ray spectroscopy (EDX)

The EDX analysis of the SCN/CD nanocomposite reveals significant insights into its elemental composition. As shown in Fig. 9(a), distinct peaks corresponding to carbon (C), nitrogen (N), oxygen (O), and sulfur (S) elements were observed at specific energy levels: the peak for carbon appears at 0.3 keV, nitrogen at 0.5 keV, oxygen at 0.8 keV, and sulfur at 2.2 keV. Quantitative analysis shows that carbon and nitrogen dominate the composition, with weight percentages of 40.5% and 56.9% respectively, suggesting their substantial incorporation into the nanocomposite structure. Oxygen contributes 2.3% by weight, while sulfur comprises a minor fraction at 0.3%. The presence



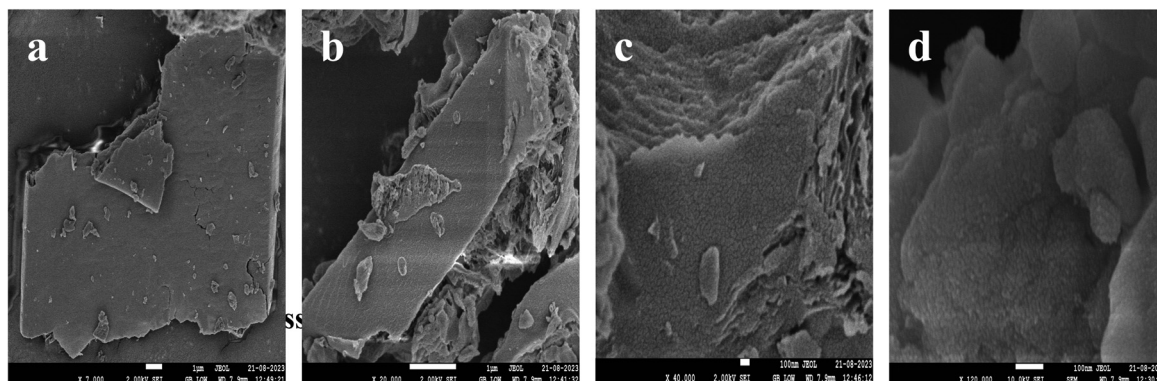


Fig. 8 SEM images of (a) SCN (b) SCN at higher magnification (c) SCN/CD (d) SCN/CD at higher magnification.

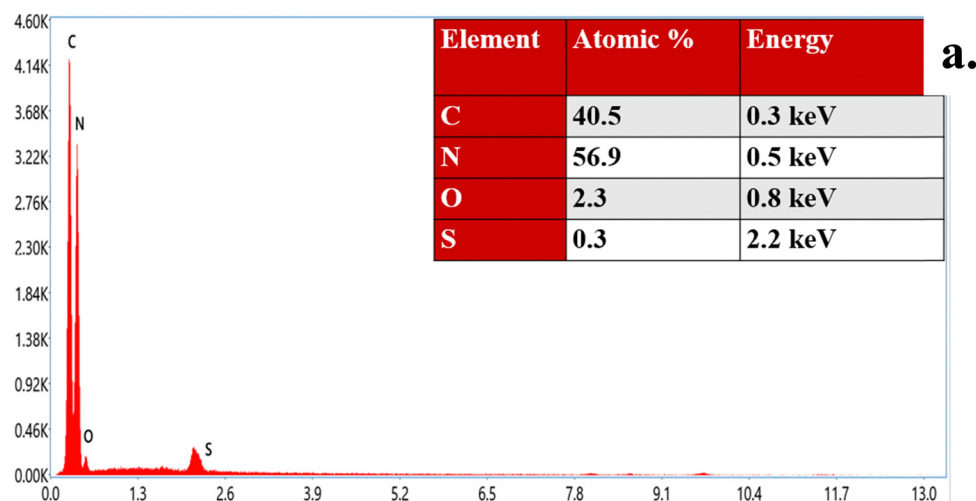


Fig. 9 (a) EDX pattern of SCN/CD.

of small amount of sulfur confirms its doping into the nanocomposite. These findings not only highlight the relative abundance of each element but also shed light on the chemical composition and potential functional groups present in the nanocomposite.

4. Results and discussion

4.1. Evaluation of catalytic performance

The photodegradation experiment was conducted within a closed wooden box, exposed to ambient air conditions, utilizing two 250-watt tungsten lamps as a visible light source. The 50 ppm solutions of organic pollutant containing TC and BPA were individually prepared in separate 1-liter volumetric flasks filled with distilled water. In the experimental setup, 50 mg of catalyst was dispersed into 100 mL of the solution containing the organic pollutants.^{44,45} To establish adsorption-desorption equilibrium, the solution was gently stirred and allowed to incubate in darkness for thirty minutes before exposure to visible light. Throughout the irradiation process, 5 mL samples of the suspension were collected at 15-minute intervals and

subsequently subjected to centrifugation to isolate the photocatalyst for analysis at specific irradiation times. The photodegradation efficiency (%) was determined using UV-visible spectrophotometry, measuring the maximum absorbance ($\lambda_{\max}$) of both compounds as shown in Fig. 10(a)–(d). We used the following equation to obtain the (%) photodegradation efficiency:⁴⁶

$$(\%) \text{ photodegradation efficiency} = \{(C_o - C_t)/C_o\} \times 100 \quad (1)$$

Assessing the photocatalytic prowess of our composite materials under visible light exposure, we employed BPA and TC as model pollutants. Both pollutants exhibited poor natural degradation properties. Among the composite materials as shown in Fig. 11(a)–(d), SCN/CD3 demonstrated the highest photocatalytic degradation rate, achieving remarkable removal rates of 92.5% for BPA and 90.7% for TC after just 60 minutes of reaction time. SCN alone also exhibited relatively effective degradation efficiencies of 68.5% for BPA and 60% for TC, attributed to its ability to absorb visible



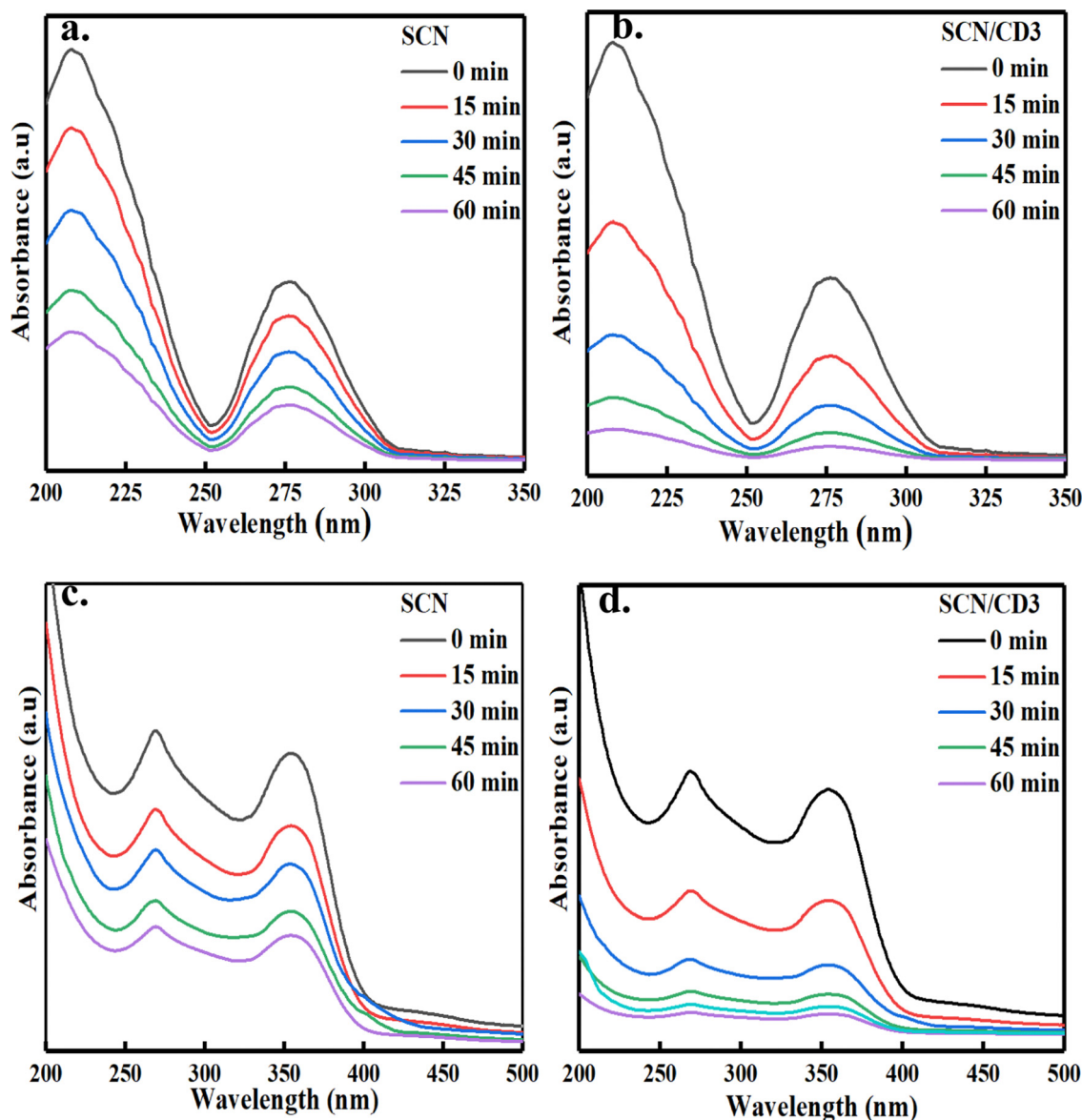


Fig. 10 UV-visible absorbance spectra of visible light degradation of BPA in presence of (a) SCN (b) SCN/CD3. UV-visible absorbance spectra of visible light degradation of TC in presence of (c) SCN (d) SCN/CD3.

light. The incorporation of CD into the composite narrowed the band gap, resulting in enhanced photosensitivity and broader absorption bands in the visible and near-infrared (NIR) light regions compared to pure SCN. This suggests that the introduction of CD acts as an electron reservoir, facilitating electron excitation in both CD and $g\text{-C}_3\text{N}_4$, thus generating more electron-hole pairs for pollutant degradation. However, an excessive concentration of CD may suppress electron-hole pair generation, underscoring the need to optimize CD concentration for optimal results. The photodegradation kinetics of BPA and TC followed pseudo-first-order kinetics. The calculated rate constants (k values) for various composite combinations against BPA and TC provide insights into their degradation efficiency as shown in Table 1. Additionally, we reviewed and compared various

studies exploring the degradation of BPA and TC using different nanomaterials, as summarized in Table 2.

4.2. Recyclability and photostability

The preservation of photocatalytic efficacy stands as a pivotal criterion in the evaluation of the durability and recyclability of the synthesized SCN/CD3 nanocomposites as photocatalysts. To gauge the potential for reuse, several degradation cycles were conducted with the SCN/CD3 nanocomposite. Following each cycle, the nanocomposite underwent centrifugation to extract it from the solution, followed by rinsing with distilled water and subsequent drying at 60 °C for two hours.^{53,54} Notably, even after undergoing five degradation cycles, the nanocomposite sustained a degradation efficiency of over 90% for BPA and maintained an efficiency of 88% for TC, as



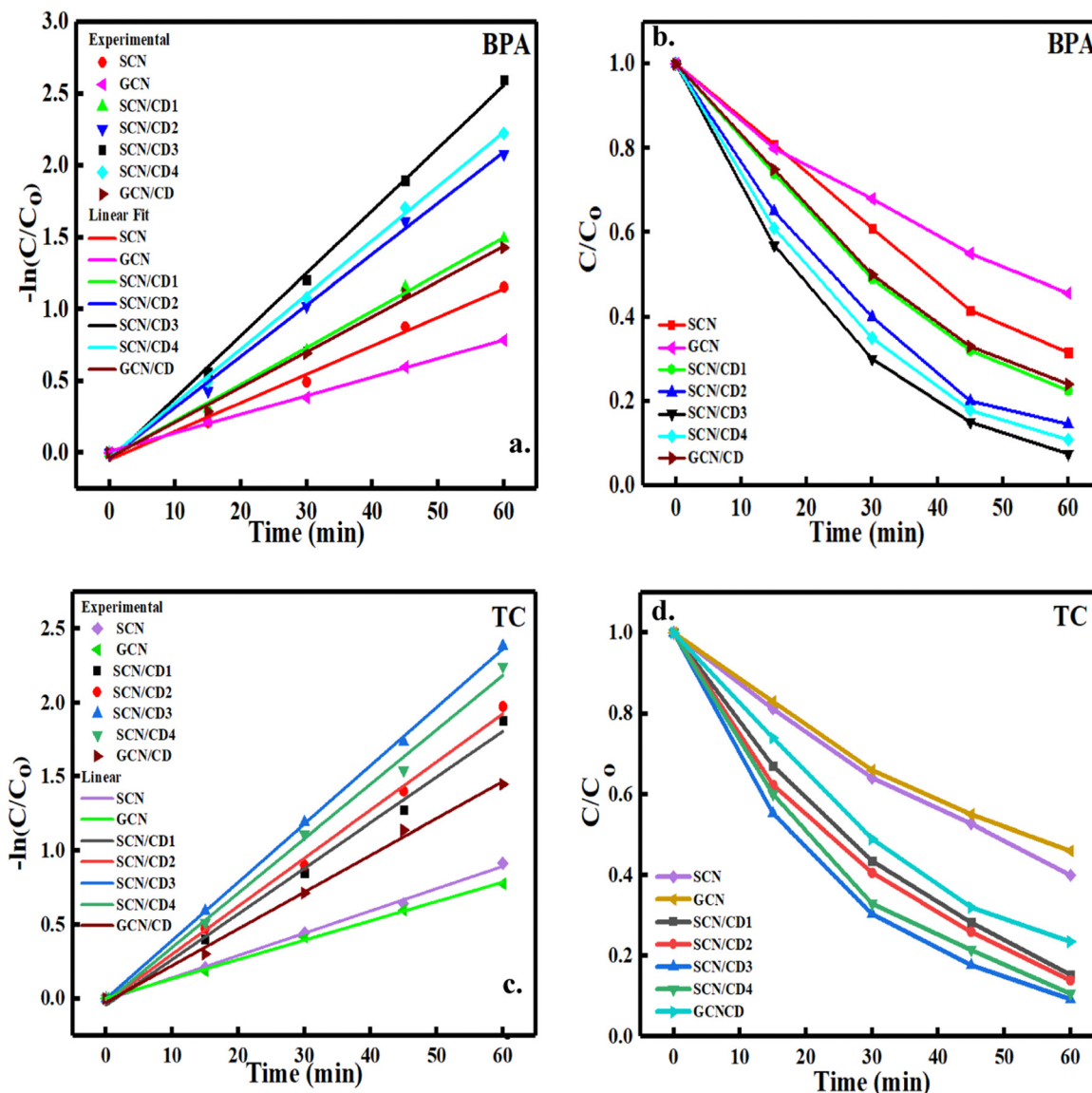


Fig. 11 (a) Photodegradation performance of BPA (b) first-order plots of BPA (c) photodegradation performance of TC (d) first-order plots of TC over the prepared nano catalysts under visible light using two 250 W Tungsten Lamp.

illustrated in Fig. 12(a). Additionally, as shown in Fig. 12(b), the XRD pattern analysis of the nanocomposite before and after degradation experiments revealed identical diffraction peaks, indicative of the robustness of the nanocomposite's structure and its enduring photocatalytic activity across multiple degradation cycles.

4.3. Radical trapping

The primary reactive species involved were determined through a free radical trapping experiment. Specifically, EDTA, isopropyl (IPA), and benzoquinone (BQ) were utilized as scavengers for holes, hydroxyl radicals, and superoxide radicals, respectively. A decrease in degradation efficiency within the reaction system typically suggests the prevalence of major active species. Each scavenger was incorporated into the reaction mixture at a concentration of 0.1 mM.⁵⁵ Comparative analysis with the absence of scavengers revealed that all scavengers exhibited a

partial suppression of photocatalytic efficiency, as depicted in Fig. 13. While IPA had a marginal effect on restraining the photodegradation efficiency of BPA and TC, the most significant reduction in the rate of photodegradation was observed in the batch treated with BQ and EDTA. This indicates that superoxide and hole radicals played pivotal roles in the photocatalytic degradation process.

4.4. Possible mechanism for degradation of BPA and TC

The degradation mechanism of BPA and TC over SCN/CD3 composite under visible light irradiation is depicted in Scheme 1, offering a nuanced understanding of the process. The determination of relative band position of SCN and SCN/CD3 was achieved through the measurement of flat band potential from Mott-Schottky plot, indicating both SCN and SCN/CD3 as an n-type semiconductor. With conduction band



Table 1 Insights into the photodegradation percentage and rate constants by different nanocomposite against BPA and TC

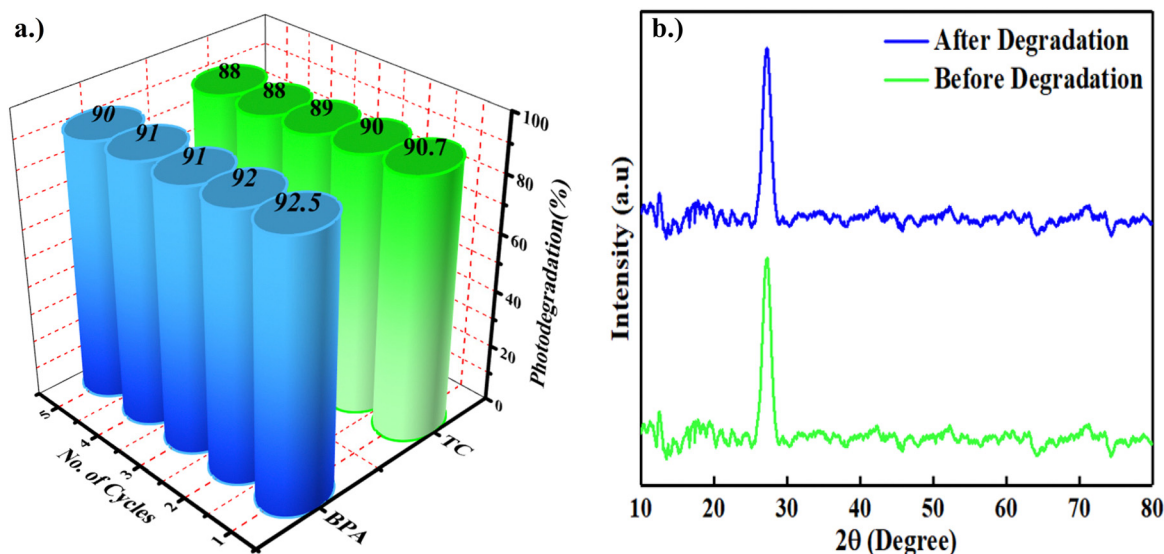
S. no.	Photocatalyst	Photodegradation efficiency (%)	Rate constant (k in min^{-1})	R^2
Performance against bisphenol A (BPA)				
1.	SCN	68.5	0.0198	0.995
2.	SCN/CD1	77.5	0.0255	0.996
3.	SCN/CD2	87.5	0.0355	0.997
4.	SCN/CD3	92.5	0.0435	0.998
5.	SCN/CD4	89.2	0.0378	0.997
6.	g-C ₃ N ₄	54.4	0.0130	0.997
7.	g-C ₃ N ₄ /CD	76.0	0.0245	0.995
Performance against tetracycline (TC)				
1.	SCN	60	0.0151	0.997
2.	SCN/CD1	84.7	0.0309	0.996
3.	SCN/CD2	86.1	0.0325	0.998
4.	SCN/CD3	90.7	0.0394	0.999
5.	SCN/CD4	89.4	0.0368	0.995
6.	g-C ₃ N ₄	54.0	0.0131	0.997
7.	g-C ₃ N ₄ /CD	76.5	0.0249	0.994

(CB) position of -1.03 eV vs. Normal hydrogen electrode (NHE) for SCN as shown in Fig. 14. By considering the optical band energy obtained from UV-DRS spectra, the band gap energy (E_g) of SCN is calculated to be 2.64 eV. Using this value, along with

the measured flat band potential, the valence band (VB) energy is calculated to be 1.61 eV vs. NHE. Similarly, for the SCN/CD3 composite, the Mott-Schottky analysis revealed the flat band potential of SCN/CD3 to be -1.57 eV, representing the energy of the conduction band. With the band gap from UV-DRS calculated as 2.60 eV, the energy of the valence band for SCN/CD3 was estimated to be 1.03 eV vs. NHE. Furthermore, the potential energy diagram vs. NHE for the composite illustrates the staggered band edge potentials between SCN and CD, forming heterojunctions at the interfaces.⁵⁶ Moreover, the CD serve a dual role, acting as both a photosensitizer and an electron mediator, thus expanding the photon absorption range effectively. Also, the incorporation of CD triggers alterations in the electron density distribution within the SCN/CD composite, thereby curbing the recombination of electron-hole pairs (e^-/h^+). Moreover, the ultra-thin structure of SCN/CD promotes swift electron transfer from the interior to the surface of the photocatalyst. Upon exposure to visible light, electrons (e^-) within the SCN/CD3 composite are excited from the VB to the CB, leading to the formation of holes (h^+) in the VB. Notably, the accumulated electrons within SCN/CD readily react with oxygen in the solution, yielding $\text{O}_2^{\bullet-}$ (0.13 eV vs. NHE). However, the system lacks the capacity to oxidize water for $\text{OH}^\bullet$

Table 2 The degradation efficiency of TC and BPA under various light irradiation compared with previous reports

S. no.	Photocatalyst	Organic pollutant	Degradation percentage (%)	Time (in min)	Light	Ref.
1.	CDs/g-C ₃ N ₄ /MoO ₃	Tetracycline	81	90	Visible	47
2.	N-CQDs/g-C ₃ N ₄ /PDS	Tetracycline	90	60	Visible	48
3.	CDs/H-CN	Tetracycline	86	120	Visible	49
4.	CDs/g-C ₃ N ₄	Bisphenol A	90	180	Visible	50
5.	RGO/g-C ₃ N ₄	Bisphenol A	90	90	Visible	51
6.	In ₂ O ₃ /OGCN	Bisphenol A	91	180	Visible	52
7.	SCN/CD	Bisphenol A	92.5	60	Visible	This work
8.	SCN/CD	Tetracycline	90.7	60	Visible	This work

**Fig. 12** (a) Five photocatalytic degradation cycles of BPA and TC using SCN/CD3 under visible light irradiation (b) the XRD patterns of the as-prepared GZP1 before and after five cycles.

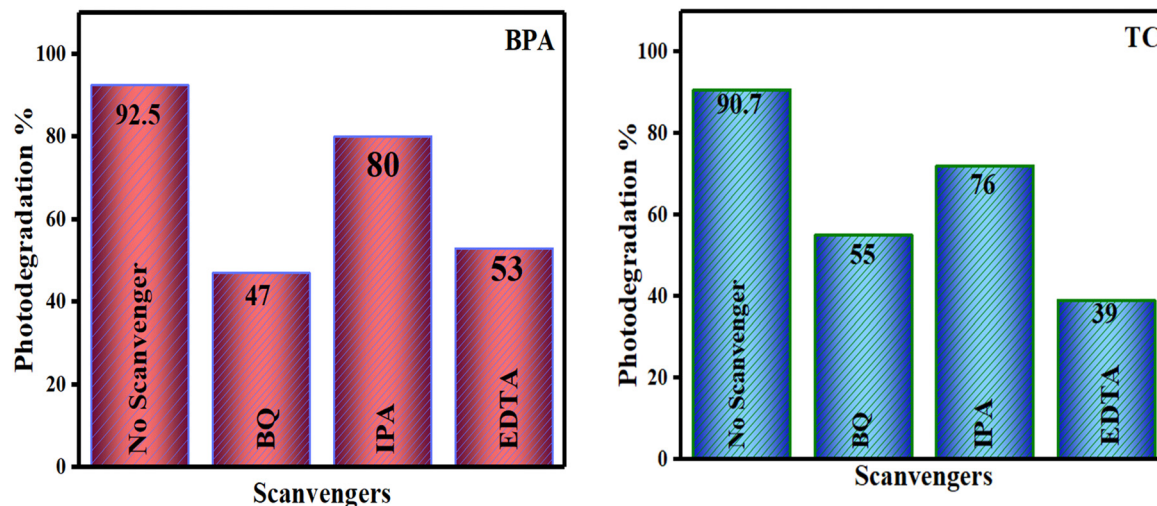


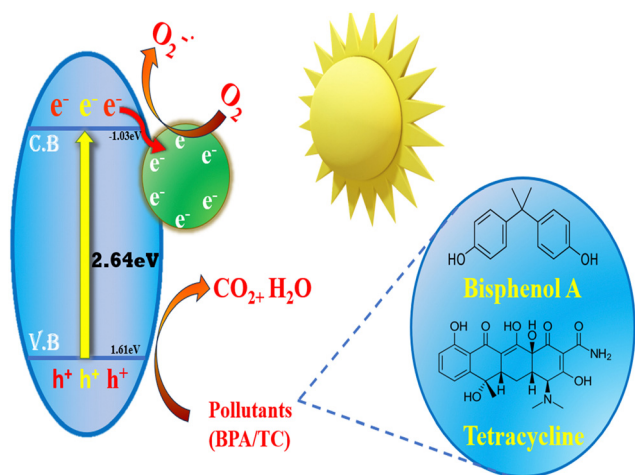
Fig. 13 Influence of radical scavengers for photodegradation of BPA and TC on SCN/CD3 under visible light irradiation.

production (2.68 vs. NHE). Consequently, $\text{OH}^\bullet$ generation occurs through the homolytic scission of H_2O_2 , originating from the reduction of $\text{O}_2^{\bullet-}$. Additionally, the holes (h^+) present in SCN/CD3 actively participate in the degradation of TC and BPA. The ultrathin structure of SCN/CD provides abundant active sites, facilitating the facile adsorption of both BPA and TC onto the photocatalyst's surface. Subsequently, these organic pollutants undergo degradation *via* the action of reactive species, notably $\text{O}_2^{\bullet-}$ and h^+ , culminating in their breakdown and complete mineralization.^{57–60} This elucidated mechanism provides a comprehensive insight into the intricate degradation processes of BPA and TC over the SCN/CD3 composite, emphasizing the pivotal roles played by CD and SCN in driving pollutant degradation under visible light irradiation.

4.5. Mechanistic degradation of BPA and TC by prepared catalyst

Liquid chromatography-mass spectrometry (LC-MS) analysis was conducted to gain insight into the intermediates formed

during the degradation process and the potential mechanistic pathway of BPA and TC in the presence of SCN/CD composites. Over a span of 60 minutes, a substantial detoxification of 92.5% for BPA and 90.7% for TC was achieved, indicating the effectiveness of the photocatalytic process. LC-MS analysis unveiled many intermediate products arising from the degradation of TC and BPA. It was observed that BPA and TC, along with their generated fragments, underwent reactions with reactive species, ultimately leading to the generation of water and carbon dioxide. For BPA, a gradual conversion into lower weight molecules such as aromatic intermediates, benzoquinones, and carboxylic acids was evident. The mechanisms underlying this degradation process involve various reactions including substitution on the aromatic ring, hydroxylation of the aromatic ring, decarboxylation, dehalogenation, bond cleavage, and ring-opening. The LC-MS analysis facilitated the identification of several intermediate products, characterized by specific mass-to-charge ratio (m/z) values. For instance, during the



Scheme 1 Schematic diagram for probable mechanism of photocatalysis.

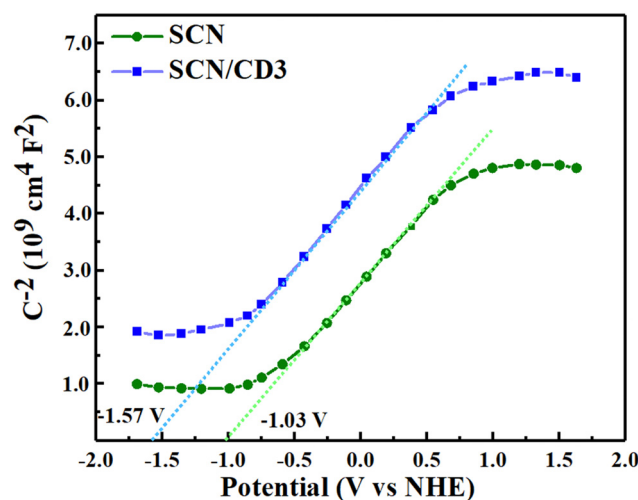


Fig. 14 Mott Schottky curve of SCN and SCN/CD3.



decomposition of BPA by the prepared photocatalyst, intermediate compounds with m/z values of 208, 197, 104, 108, 152, 122, and 135 were detected.⁶¹ Similarly, active species targeted double bonds, aromatic rings, amino groups, leading to ring-opening reactions, loss of functional groups, or rupture of primary carbon bonds, thereby generating many intermediates in the case of TC. The intermediate compounds formed during the decomposition of TC exhibited m/z values of 445, 272, 238, 433, 301, 149, 210, 438, 139, and 103.⁶² This comprehensive LC-MS analysis provides valuable insights into the intermediates formed during the degradation process, shedding light on the potential degradation pathways followed by BPA and TC under photocatalytic conditions. Information regarding the proposed degradation pathway and intermediates is provided in S1 (ESI[†]), offering a detailed elucidation of the mechanisms underlying the photocatalytic degradation of BPA and TC.

5. Conclusion

The sulfur modification of g-C₃N₄ coupled with the integration of CD into the nanocomposite yields a significant enhancement in visible light activity and efficiency for photocatalytic degradation of CECs like BPA and TC. Within the SCN material, the presence of CD serves as electron reservoirs, effectively mitigating electron-hole recombination and thereby augmenting pollutant degradation. Particularly noteworthy is the exceptional efficiency demonstrated by the SCN/CD3 photocatalyst, achieving BPA degradation rates of up to 92.6% and TC degradation rates of up to 90.7% within a brief 60-minute irradiation period under visible light. Remarkably, the catalyst maintains high degradation efficiency over five cycles, with superoxide radicals (O₂^{•−}) and holes (h⁺) identified as the primary reactive species responsible for pollutant degradation. Broadening the scope of pollutant targets beyond BPA and TC to encompass a wider range of CECs prevalent in various environmental matrices will increase the applicability and relevance of SCN/CD composites in environmental remediation efforts. These endeavours hold significant promise for making substantial contributions to sustainable pollution mitigation strategies.

Author contributions

Ankoo Sura: conceptualization, formal analysis, investigation, writing – original draft preparation, methodology, visualization. Arjun Singh, Sudha Narwal, Priya Malik and Amanvir Singh: data curation, formal analysis, visualization and investigation. Sonia Nain and Surender Duhan: project administration, conceptualization, formal analysis, resources, investigation, supervision, reviewing and editing.

Conflicts of interest

The authors declare no competing financial and/or non-financial interests.

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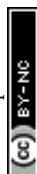
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