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Cobalt doped titania-carbon nanosheets with induced oxygen vacancies for photocatalytic degradation of uranium complexes in radioactive wastes[†]

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The photocatalytic degradation of uranium complexes is considered among the most efficient techniques for the efficient removal of uranium ions/complexes from radioactive wastewater. Described here is a nanostructured photocatalyst based on a cobalt-doped TiO_2 composite with induced oxygen vacancies ($\text{Co@TiO}_2\text{-C}$) for the photocatalytic removal of uranium complexes from contaminated water. The synergy between oxygen vacancies and Co-doping produced a material with a 1.7 eV bandgap, while the carbon network facilitates electron movement and hinders the e-h recombination. As a result, the new photocatalyst enables the decomposition of uranium-arsenazo III complexes (U-ARZ3), followed by photocatalytic reduction of hexavalent uranium to insoluble tetravalent uranium. Combined with the nanosheet structure's high surface area, the photocatalytic decomposition, reduction efficiency, and kinetics were significantly enhanced, achieving almost complete U(VI) removal in less than 20 minutes from solution with a concentration as high as 1000 mL g^{-1} . Moreover, the designed photocatalyst exhibits excellent stability and reusability without decreasing the photocatalytic performance after 5 cycles.

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1. Introduction

The use of nuclear-powered electricity for diagnosing and treating diseases and other purposes has increased the possibility of radioactive species being leaked into the environment.^{1,2} Due to its radiochemical and toxicological effects, particular concern has been given to uranium, among other radioactive waste components.³ Uranium forms complexes in waste solutions with existing chelating agents such as tributyl phosphate, Alizarin red S, ethylenediaminetetraacetic acid (EDTA), xylanol orange, arsenazo III, and chlorophosphonazo.^{4–6} Such complexes are absorbed into the human body until they reach the liver and kidneys. Several methods have been used or

suggested to remove uranium and other heavy metal ions from a nuclear waste solution, such as chemical precipitation, adsorption, membrane filtration, and ion exchange.^{7–14} However, photocatalytic degradation is receiving increasing interest for removing radionuclide complexes from contaminated solutions due to the low cost of operation, simplicity, and environmental friendliness.^{15–19} Recent work has revealed that photo-generated electrons can convert uranium into an insoluble form to reduce the soluble hexavalent state to its insoluble quaternary state.^{20,21}

Titanium-based nanostructures are commonly used as photocatalysts to remove a wide range of organic and inorganic compounds. Coupled with its abundance, the excellent chemical and thermal stability of TiO_2 make it one of the best candidates for photocatalysts.²² TiO_2 exists in three different polymorphs (*i.e.*, anatase, rutile, and brookite). These polymorphs exhibit different physical characteristics, such as the differences in their bandgap energies (*i.e.*, 3.0 eV for rutile and 3.2 eV for both anatase and brookite).²³ The photocatalysis mechanism exerted by TiO_2 can be carried out by forming free electrons and holes in the conduction and valence band region.^{16,22}

Uranyl ion reduction on TiO_2 by ultraviolet (UV) irradiation was reported for the first time by Amadelli *et al.*²⁴ Several studies

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have since looked at the photocatalytic reduction of U(vi) on TiO₂ photocatalysts.^{25–49} Kim *et al.* demonstrated that TiO₂ electrodes could photoelectrochemically reduce and eliminate U(vi) spikes in natural seawater.³⁵ Li *et al.* also investigated TiO₂ photocatalysis in uranium recovery from seawater. The photocatalytic approach demonstrated high efficiency for reducing and extracting U(vi) under different circumstances, with the reduced uranium deposits on TiO₂ easily recovered by using Na₂CO₃ solution.⁴⁶ Li *et al.* used TiO₂/Fe₃O₄ and TiO₂/graphene/Fe₃O₄ composites as photocatalysts to boost the photocatalytic activity of TiO₂ for U(vi) reduction.⁴¹ The photo-generated e[−] in the TiO₂ conduction band (CB) was partially transferred to the Fe₃O₄ surface, reducing Fe(II) and Fe(III) in the structural Fe₃O₄ phases when exposed to UV light. Therefore, U(vi) ions are reduced by both the generated Fe species and the photoelectrons, significantly enhancing the U(vi) removal. Although Li *et al.* reported that the photo-reactivity decreased when graphene was added, the graphene layer inhibits the photo-dissolution of Fe₃O₄, making the ternary composite considerably more stable. Therefore, designing a stable photocatalyst that can efficiently reduce U(vi) without contaminating the wastewater with the reaction by-product is essential for the wider application of photocatalytic treatment of nuclear waste. More importantly, most of the studies published so far have considered standard U(vi) solutions and ignored the organic matter that commonly exists in real waste solutions. These organic species influence the photocatalytic reduction of U(vi) by two mechanisms; (a) the adsorption of U(vi) on solid surfaces is altered by the competing diverse organics,^{50,51} and (b) U(vi) forms complex ions with a lot of this organic matter, which completely changes the nature of the uranium ions in the solutions.^{4–6}

The present study introduces a nanostructured TiO₂-based photocatalyst (Co@TiO₂-C) to efficiently remove uranium complexes from contaminated water. Co@TiO₂-C promotes two photocatalytic processes: decomposing the complexes, such as U(vi)-ARZ3, and reducing U(vi) to insoluble U(IV). We carefully designed porous nanostructures of connected 2D nanosheets of cobalt-doped anatase with induced oxygen vacancies. The synergy between the oxygen vacancies and the cobalt distortion results in a material with a low bandgap of 1.7 eV, facilitating photoelectron transfer. The oxygen vacancies were introduced *via* partial carbothermal reduction during nanophotocatalyst preparation. The uranium complex removal mechanism was also studied using several *ex situ* spectroscopic techniques. A satisfactory U(vi) removal can be achieved even at low concentrations. Notably, the photoreduction efficiency of 99.6% remains achievable by U(vi)-ARZ3 to form (UO₂O₂·2H₂O). The photocatalysis method using Co@TiO₂-C was demonstrated to be a promising approach for complex radioactive waste remediation.

2. Materials and methods

2.1. Chemicals

Titanium isopropoxide Ti[OCH(CH₃)₂]₄, cobalt acetate (CH₃CO₂)₂Co, potassium chloride (KCl), ethanol (C₂H₅OH), glucose

(C₆H₁₂O₆), arsenazo III ((HO)₂C₁₀H₂(SO₃H)₂(N=NC₆H₄AsO₃−H₂)₂), and uranyl nitrate UO₂(NO₃)₂·6H₂O were Sigma-Aldrich products (Germany). Deionized water (DIW) used in all experimental steps has an electric conductivity of 0.7 μS cm^{−1}.

2.2. Synthesis of the Co/C doped TiO₂ photocatalyst

About 0.5 mL of aqueous KCl (0.1 M), 0.88 g of cobalt acetate, and 1.5 mL of titanium isopropoxide were added to 100 mL ethanol and stirred for 4 h. The solution was then transferred to a 100 mL stainless steel autoclave lined with Teflon and heated at 170 °C for 30 h before being air-cooled to room temperature. The resultant product was collected and thoroughly cleaned with ethanol before being dried in an oven at 60 °C. To obtain Co@TiO₂, the dried powder was sintered at 500 °C for 3 h. To prepare Co@TiO₂-C, 0.42 g of the obtained Co@TiO₂ composite was dispersed in 75 mL of DIW containing 0.019 g glucose. The liquid was stirred for 2 h before being transferred to a 100 mL Teflon-lined stainless steel autoclave. The autoclave was kept at 170 °C for 20 h before being air-cooled to ambient temperature. The resultant product was collected and thoroughly washed with water several times before being calcined in an oven at 500 °C for 3 h to generate an amorphous Co@TiO₂-C photocatalyst.

2.3. Photocatalyst performance

Photodegradation/reduction of the U(vi)-ARZ3 complex and free ARZ3 and U(vi) in an aqueous solution was carried out under UV irradiation. The batch experiments were conducted in a photoreactor at 25 ± 1 °C under ambient conditions. 10 mg of the photocatalyst was added into 10 mL of 100 mg L^{−1} of U(IV)-ARZ3 (S/L ratio 1:1) at different pH values. A similar experiment was carried out without ARZ3 for comparison. After a certain time, an aliquot (5 mL) of the solution was taken out and filtered for analysis. The concentration of U(vi) was determined using ICP. The degradation of U(vi)-ARZ3 complexes was determined at 450 and 650 nm using UV-vis spectrometry. The removal rate of all samples was calculated using the equation: $R\% = [(C_0 - C_t)/C_0] \times 100$; where C_0 is the initial concentration and C_t is the concentration at time t .

3. Results and discussion

3.1. Characterization of the Co@TiO₂-C photocatalyst

The phase characterization of the as-obtained Co@TiO₂-C nanostructure was first determined by the XRD technique. The XRD pattern for the Co@TiO₂-C nanostructure is presented in Fig. 1a. In the XRD pattern, the characteristic diffraction peaks (101), (004), (200), (105), (211), (204), (116), (220) and (215) of pure TiO₂ can be detected. The peak for the 101 plane slightly shifts from $2\theta = 25.28^\circ$ for pure anatase *vs.* $2\theta = 25.44^\circ$ for Co@TiO₂-C.⁵² This shift is caused by changes in the local structure around Ti⁴⁺ following Co²⁺ substitution and oxygen vacancy creation.⁵³ The crystal size of the Co@TiO₂-C nanocomposite sample is calculated using Scherrer's equation to be 11.38 nm. It should be noted that the XRD spectrum of the composite showed some minor peaks of sub-oxides and



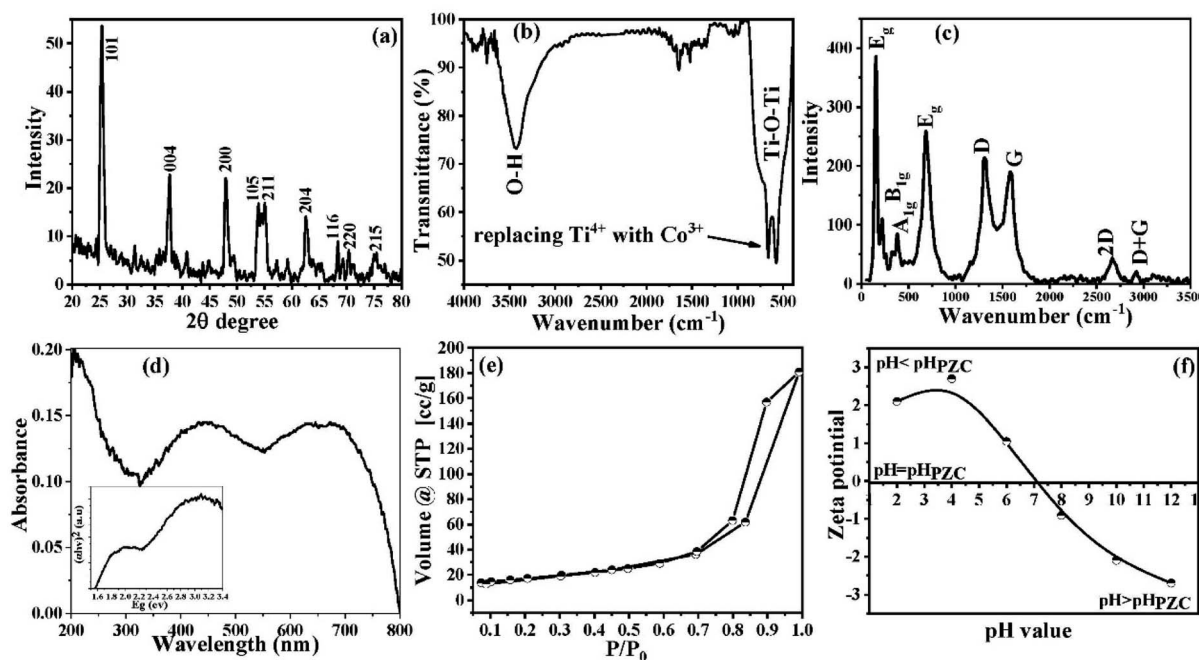


Fig. 1 (a) XRD patterns, (b) FTIR spectrum, (c) Raman spectrum, (d) UV-vis spectroscopy, (d inset) the energy bandgap, (e) N₂ adsorption/desorption isotherm, and (f) zeta potential in 0.01 M KCl as a function of pH of the synthesized Co@TiO₂-C photocatalyst.

Magneli phases (the unlabeled peaks in Fig. 1a), which can be attributed to the partial reduction of the oxides. The percentage of the sub-oxides and Magneli phases was calculated to be around 9%. It is well known that these phases are conductive and therefore increase the overall conductivity of the composite.^{54,55} We believe that these nonstoichiometric phases have contributed to photocatalytic activities.

The surface chemistry of the Co@TiO₂-C photocatalyst is investigated using the FTIR spectrum (Fig. 1b). The FTIR spectrum revealed a high-intensity vibration band at 3425 cm⁻¹ attributed to the O-H group's stretching vibration.⁵⁶ The vibration peak for the adsorbed water molecules on the surface of the nanoparticles is at 1643 cm⁻¹ (bending vibration mode). The vibration band at ~1396 cm⁻¹ is associated with the bending vibration of the C-H bond. High-intensity deformation bands between 400 and 800 cm⁻¹ correspond to the Ti-O-Ti vibrational mode.^{57,58} This deformation is associated with replacing Ti⁴⁺ with Co³⁺ within the anatase network.

The Raman spectrum acquired for the Co@TiO₂-C sample is illustrated in Fig. 1c. Peaks at 151, 390, 510, and 627 cm⁻¹ are ascribed to anatase phase modes E_g, B_{1g}, A_{1g}, and E_g, respectively.⁵⁹ The strength of the B_{1g}, A_{1g}, and E_g peaks decreased after Co doping and carbothermal reduction (Fig. S1a†). This result suggests that the Co did not entirely transform the anatase phase and implies considerable deformation in the long-order crystallite, consistent with the XRD and FTIR results. The detection peaks at the D and G band locations (1300 cm⁻¹ and 1850 cm⁻¹, respectively) demonstrates the existence of carbon despite the calcination in air with Ti-Co oxides.^{60,61} The peak intensity ratio I_D/I_G is 1.33, indicating that the carbon is most likely amorphous.

UV-vis spectroscopy confirms the substitution of Ti⁴⁺ with the Co dopant. The absorption peak of pure anatase at 363 nm corresponds to the intrinsic band-band transition.⁶² The Co@TiO₂-C spectrum shows two peaks at 495 and 582 nm (Fig. 1d). The peak at a lower wavelength is associated with the ⁶A_{1g}(S) to ⁴A_{1g}(G), ⁴E_g(G), and A_{1g}(S) to ⁴T_{2g}(G) transitions, whereas the peak at 582 nm is for the Co²⁺ in octahedral coordination ⁶A_{1g}(S) to ⁴T_{1g}(G). Each Ti⁴⁺ in TiO₂ is coordinated with six oxygen atoms. When Co²⁺ substitutes Ti⁴⁺, the electrons in the d-orbital of Co²⁺ experience repulsion, resulting in the splitting of Co²⁺ d-orbitals, showing that Co²⁺ is inside the lattice.

The energy band gap of Co@TiO₂-C can be determined from the linear portion of the Tauc plot, which was fitted and extrapolated for zero absorbance, *i.e.*, $F[R] = 0$.⁶³ The energy bandgap of Co@TiO₂-C is 1.6 eV decreased from 3.2 eV in pure anatase (inset of Fig. 1d). Such energy levels indicate a shift in the local distribution of Ti⁴⁺ or partial conversion of Ti⁴⁺ to Ti³⁺ and the formation of empty oxygen sites. The change in the bandgap of Co@TiO₂-C can be explained by the reduced particle size and the synergy between the Co substitution and the oxygen vacancies caused by the regulated carbothermal process.

The N₂ adsorption/desorption technique was used to investigate the porosity structure of the Co@TiO₂-C nanocomposite. As shown in Fig. 1e, the Co@TiO₂-C isotherm exhibits typical characteristics for type IV isotherms, according to the IUPAC classification.⁶⁴ At $P/P_0 \geq 0.7$, the isotherm displays a distinguished hysteresis, indicating a large number of mesopores (Fig. 1e). The isotherm also exhibits an increasing trend at low pressure, corresponding to adsorption at micropores. Because of the hierarchical porous structure with nanochannels between



the oxide particles, it is possible to deduce that there is a multilayer range of physisorption isotherms. Co@TiO₂-C has an estimated S_{BET} of 60.946 m² g⁻¹. As a result of co-doping and partial carbothermal reduction, certain new pores are introduced into the structure. One explanation for the increase in S_{BET} might be that oxygen or CO_x gases are exiting the oxide structures, which relates to creating oxygen vacancies. Furthermore, the nanoparticles are arranged into 2D nanosheets, as will be discussed with the SEM images, reducing aggregation and creating additional surfaces. The point of zero charges (pzc) of the synthesized Co@TiO₂-C photocatalyst was found to be 7.1 (Fig. 1f). This high pH_{PZC} value resulted from cobalt and oxygen vacancies, and it is virtually consistent with the FTIR spectrum data since it verifies the increase in the absorption of the hydroxyl group at 3425 cm⁻¹.⁶⁵

The SEM photos clearly show the morphology of the Co@TiO₂-C nanostructure (Fig. 2a). The morphology has a definite 2D structure, indicating that TiO₂ nanoparticles tend to cluster in the form of sheets. The nanosheet structure is maintained following cobalt additions, and partial carbothermal reduction implies preferential development directions. The 2D nature of the morphology is most likely due to the increase in the density of states perpendicular to the C-axes caused by Co doping of the Ti site and the partial elimination of oxygen by carbon. The porous zig-zag structure is formed by loosely linking the 2D sheets together. More information may be gleaned from the TEM pictures (Fig. 2b), which revealed that the material is made up of nanoparticles with parallel facets and

an average size of about 15 nm that shows a uniform distribution with pore ordering over graphite sheets (Fig. 2c and d).

XPS was performed on Co@TiO₂-C at room temperature to further estimate the possibility of secondary phases, oxygen deficiency, and oxidation states of Ti and Co in the near-surface region, as shown in Fig. 3. The peaks in the survey (Fig. 3a) at 459.17 eV, 781.5 eV, 530.8 eV, and 285.82 eV can be attributed to Ti 2p, Co 2p, O 1s, and C 1s elements, respectively.⁶⁶⁻⁶⁸ The high-resolution Ti 2p spectrum is shown in Fig. 3b. The presence of a pair of symmetric spin-orbit doublets with binding energies corresponding to Ti 2p_{3/2} and Ti 2p_{1/2} for the composite is verified by two large peaks at 458.24 eV and 463.99 eV, corresponding to Ti⁴⁺ in the TiO₂ phase.⁶⁶⁻⁶⁹ Some peaks evidence the existence of the Ti³⁺ and Ti²⁺ oxidation states, together with Ti⁴⁺, proving the Co-doping and the partial reduction in the anatase crystals. The peaks at 457.78 eV and 460.25 eV correspond to Ti 2p_{3/2} and Ti 2p_{1/2} of Ti³⁺.^{69,70} Also, the peaks at 458.76 eV and 464.07 eV are for Ti 2p_{3/2} and Ti 2p_{1/2} of Ti²⁺.^{71,72} More evidence of doping can be obtained from the Co 2p spectrum (Fig. 3c). The spectrum is best fitted to two spin-orbit doublets characteristic of Co²⁺ and Co³⁺, where two peaks of Co 2p_{3/2} and Co 2p_{1/2} are located at 780.01 eV (Co³⁺) and 796.01 eV (Co²⁺). The two conspicuous shake-up satellite peaks are located at 785.92 eV and 803.22 eV, revealing the co-existence of Co²⁺ and Co³⁺. The other less intense peaks that appeared for Co 2p_{3/2} and Co 2p_{1/2} are located at 781.98 eV (Co²⁺) and 799.64 eV (Co³⁺).⁷² The co-existence of different states of cobalt suggests that carbothermal partial reduction has also taken place in the

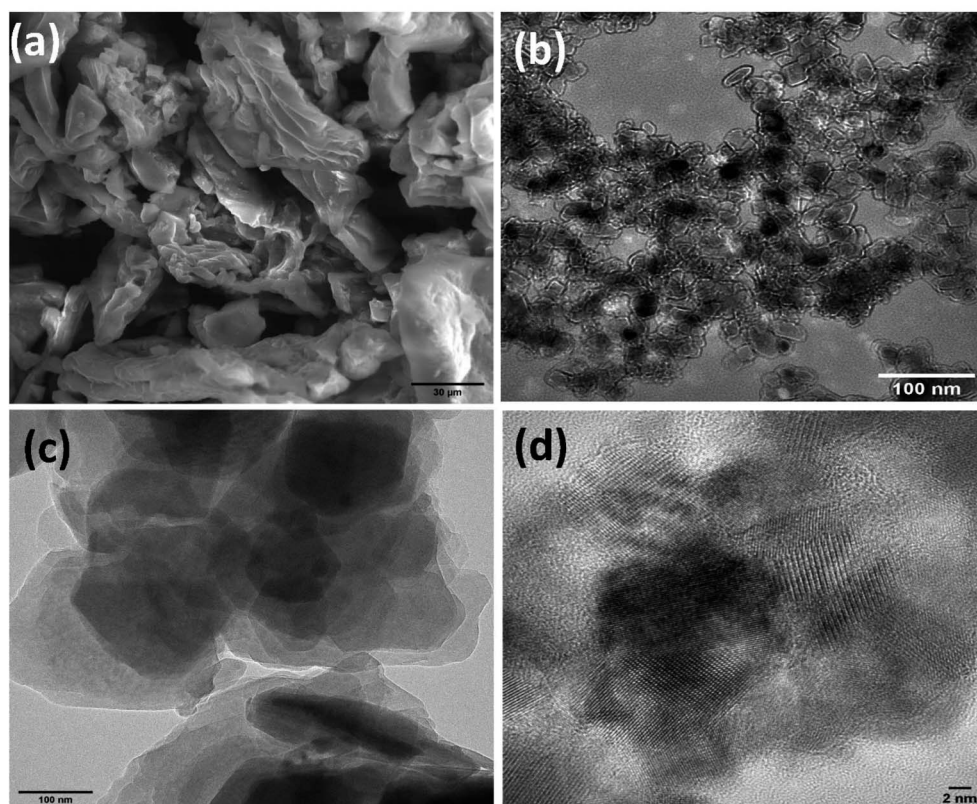


Fig. 2 SEM (a) image and TEM image (b–d) of the synthesized Co@TiO₂-C photocatalyst.



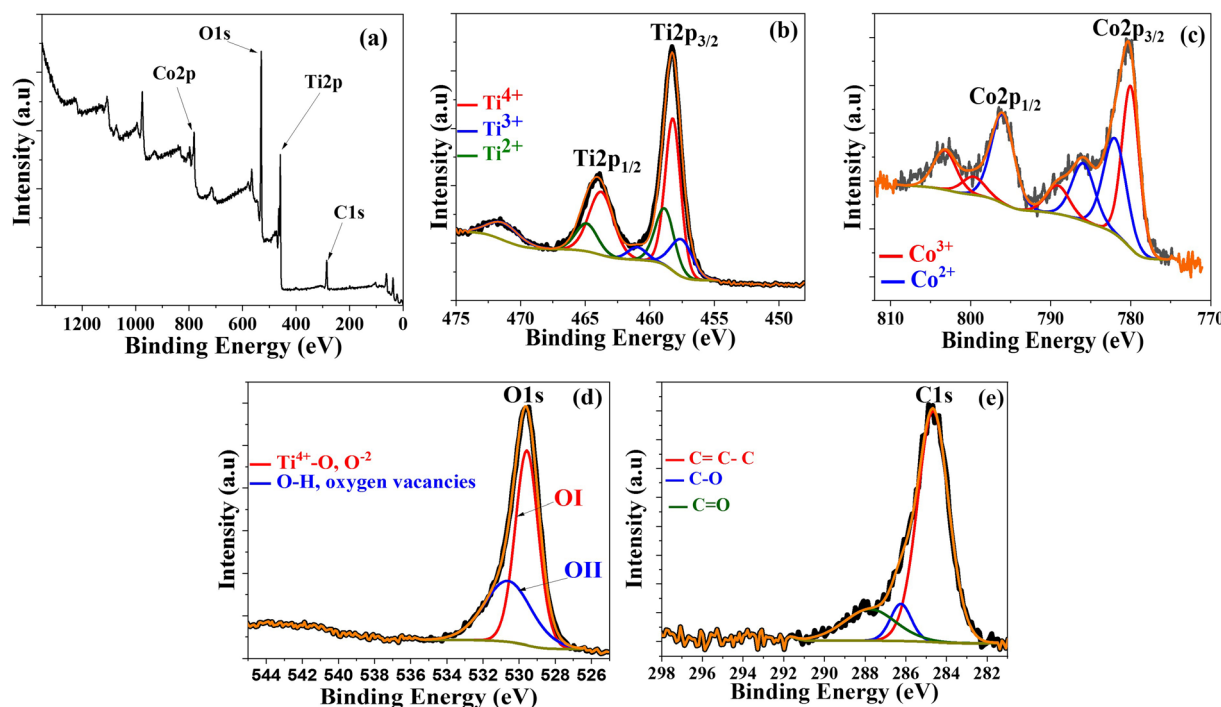


Fig. 3 XPS spectra of Co@TiO₂-C, (a) Survey spectrum, (b) Ti 2p, (c) Co 2p, (d) O 1s and (e) C 1s core-levels of the Co@TiO₂-C.

cobalt sites in the crystals. The high-resolution O 1s spectra (Fig. 3d) display two peaks at 529.57 eV and 530.62 eV. The low energy peak is attributed to O atoms bonded to metals (Ti⁴⁺-O), whereas the high-energy one can be attributed to oxygen vacancies or surface contamination by hydroxyl species.⁶⁹⁻⁷⁴ The C 1s spectra revealed three distinct peaks at 284.66, 286.24, and 287.76 eV (Fig. 3e). The first characteristic peak, at 284.66 eV, is for the C-C bond of the graphitic carbon on the surface of Co/TiO₂. The other two peaks, at 286.24 and 287.76 eV, are for the C-O-C and C=O bonds, suggesting that the carbon surface layer is bonded to Co@TiO₂ *via* oxide functional groups.⁶⁶

3.2. Photocatalytic performance of Co@TiO₂-C catalyst

Uranium-ARZ3 complex degradation/reduction experiments using the Co@TiO₂-C photocatalyst were carried out under UV irradiation. First, we tested the adsorption capability of Co@TiO₂-C by stirring uranium nitrate or the U(vi)-ARZ3 complex with a concentration of 0.1 mM in the dark. The synthesized catalyst showed weak U(vi) adsorption in the dark (Fig. 4a). We then tested the photo-stability of the uranium nitrate, ARZ3, and U(vi)-ARZ3 complex solutions by exposing them to UV light without a photocatalyst. No noticeable changes in the concentrations can be observed, indicating good photo-stability (Fig. 4b). It is only when the solutions are irradiated with UV light in the presence of Co@TiO₂-C that we start to observe a noticeable reduction in the uranium concentrations. The decrease in the peak intensities at $\lambda_{\text{max}} = 450$ nm and $\lambda_{\text{max}} = 650$ nm in the ARZ3 dye and U(vi)-ARZ3 complex, respectively, can be attributed to the breaking of the bond N=N present in their chemical structure (Fig. 4c).⁵⁰ For the U(vi)-ARZ3 solution,

the N-N bond breaking is also associated with a decrease in the uranium concentration. These results imply a reduction in soluble hexavalent uranium U(vi) to insoluble tetravalent uranium U(iv) *via* a two-step photocatalytic process.¹⁸ Supplementary to this, U(iv) nitrate in water/ethanol (10% V/V) was used to evaluate the photocatalytic activity of the synthesized photocatalyst towards fee U(iv) ions. The concentration of U(vi) decreased sharply in the first few minutes after irradiation, showing that the photocatalysis process was crucial for the conversion of U(vi) to an insoluble product (U(iv)) (Fig. 4d).

It is well known that the pH value strongly affects the photo-reaction process. For that reason, we studied the effect of pH on the removal of uranium from different solutions. As seen from Fig. 5a, all solutions' degradation occurs at pH values from 2 to 8. The removal efficiency achieves peak values at almost the same pH value, which is very close to the measured pH_{PZC} of the solid surface of the Co@TiO₂-C catalyst (about 7.1).

Without uranium in the solution, the highest rate of ARZ3 dye degradation was 98.1% at pH 4 (Fig. 5a). It should be mentioned that ARZ3 exists as an anion and gets a negative charge at low pH. The surface of Co@TiO₂-C has positive charges in acidic media, resulting in repulsion forces with anionic ARZ3, and this hinders the degradation process.⁷⁵ The difference in the symmetry of the ARZ3 bonds in acidic and basic media plays an important role in the degradation process. ARZ3 is found in the azo-form, and ionization of the acidic (-OH and -NH-N) or basic (-N₂⁻ and CO) groups leads to symmetry disruption and the appearance of quinone-hydrazone forms together with the azide.⁷⁶ Fig. S1b† shows the structure of the arsenazo III and uranium/arsenazo III complex.



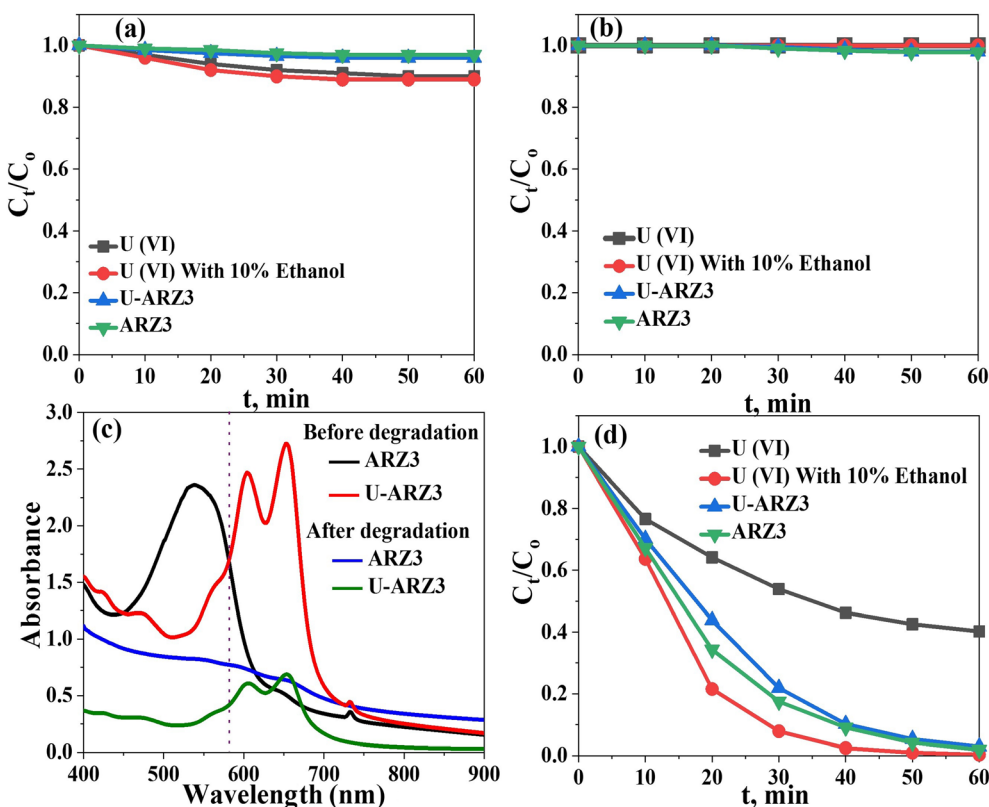


Fig. 4 (a) Effect of UV light on U(VI) and ARZ3 solutions without Co@TiO₂-C catalyst. (b) Effect of Co@TiO₂-C in the dark. (c) Effect of Co@TiO₂-C in UV light. (d) Absorbance spectra of ARZ3 and the U(VI)-ARZ3 complex.

The best degradation efficiency of U(VI)-ARZ3 is about 97.1%. The best efficiency in aqueous solutions and the water/ethanol mixture is 59.8% and 99.69%, respectively. The peak in the photo-reduction efficiency of Co@TiO₂-C at nearly the pH_{PZC} is due to the presence of different uranium species, as displayed by Visual MINTEQ application (Fig. 5b). As a result of the forces of attraction and electrostatic repulsion between these species and the surface charge of Co@TiO₂-C, protons are generated on the surface and hinder further adsorption of U(VI)

ions (*i.e.*, mainly with UO₂²⁺). These protons also compete with the positive uranium ions to consume the photo-generated electron e⁻ and thus reduce the photo-reaction activity. At pH \geq 6, U(VI) exists in the form of neutrally and negatively charged ions (mainly UO₂(OH)₂ and UO₂(OH)₃⁻) which also generate electrostatic repulsion between them and the negative charges of the surface of Co@TiO₂-C. As such, the photocatalytic activity also decreases at high pH.^{29,77-79} The low removal of uranium ions from the U(VI)-ARZ3 solution is likely due to the facile

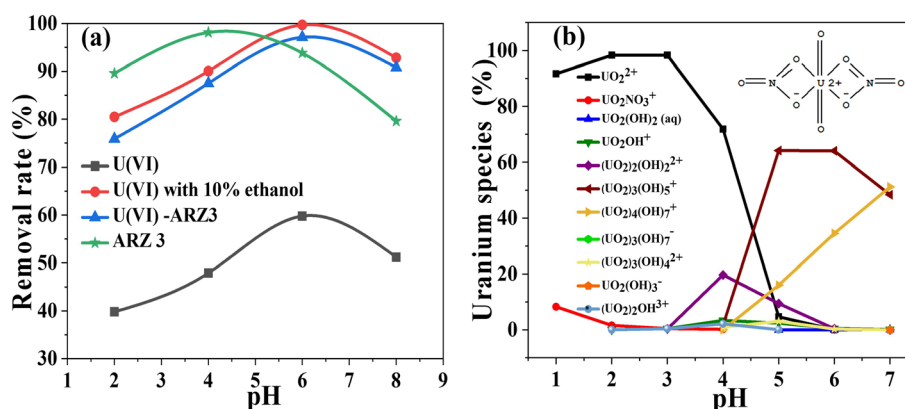


Fig. 5 (a) The effect of pH on the degradation/reduction of 50 mg L⁻¹ U(VI)-ARZ3, free ARZ3 dye, and free U(VI) ions using the Co@TiO₂-C photocatalyst for 60 min (solution volume; 10 mL, catalyst weight; 10 mg) and (b) speciation of uranium in aqueous solution at different pH values.

adsorption of the ARZ3 anions on the positively charged surface catalyst, leaving UO_2^{+2} in the solution. On the other hand, the variation in photo-reduction efficiencies of free U(vi) ions in water compared to in the water/ethanol mixture is due to the ability of ethanol to scavenge the photo-hole h^+ which are generated in the valence band (VB). These holes could re-oxidize U(iv) back to soluble U(vi).¹⁸ The re-oxidation of U(iv) into U(vi) could also take place by superoxide radical (*i.e.*, $\cdot\text{O}_2^-$) radicals, which are derived from the dissolved atmospheric O_2 . The concentration of these radicals significantly reduces when 10% ethanol is added to the solution.

To shed more light on the kinetics of photodegradation/reduction of U(vi)-ARZ3, we first monitored the changes in absorption spectra of the degradation of the $[\text{U-ARZ3}]^{n+}$ complex after the addition of the Co@TiO₂-C catalyst as a function of time (Fig. S2†). Within the first 10 min, the amount of U(vi)-ARZ3 removed using the Co@TiO₂-C catalyst is 29.8%, and equilibrium is acquired within 60 min. The photocatalytic removal data were applied to a pseudo-first-order model: $\ln(C_t/C_0) = -Kt$, where C_0 and C_t are the initial and final concentrations after time t of exposure to UV radiation, and K is the rate constant. The kinetic parameter was determined from the slope of plotting $\ln(C_t/C_0)$ versus t (Fig. 6a-d). The regression coefficient (R^2) values for the removal of U(vi)-ARZ3, free ARZ3, and U(vi) are calculated to be > 0.98 (Table S1†), indicating that the photodegradation/reduction processes can be

well-fitted with the pseudo-first-order kinetic. At the optimum pH, Co@TiO₂-C had the greatest apparent reaction rate constants of $6.155 \times 10^{-2} \text{ min}^{-1}$, $6.781 \times 10^{-2} \text{ min}^{-1}$, $9.97 \times 10^{-2} \text{ min}^{-1}$ and $1.514 \times 10^{-2} \text{ min}^{-1}$ for the U(vi)-ARZ3 complex, free ARZ3 dye, free U(vi) in water/ethanol and free U(vi) in aqueous media, respectively.

The initial concentration of the solution is an important factor in determining the performance of the photocatalysts. Therefore, the removal of various ions is investigated in solutions with concentrations ranging between 1 and 1000 mg L⁻¹ under the optimal experimental conditions (*i.e.*, pH 6, contact time; 60 min, S/L; 1 g L⁻¹, and temp. 25 °C). As shown in (Fig. 7a-d), Co@TiO₂-C appeared to remove the free ARZ3 dye at concentrations $\leq 100 \text{ mg L}^{-1}$ with an efficiency of up to 97%. By increasing the concentrations, the efficiency of the photocatalyst increases and then gradually decreases until it reaches 61% at a concentration of 1000 mg L⁻¹. For free U(vi) in pure water, the photocatalytic process cannot fully remove the uranium ions, even at a very low concentration. A significant improvement could be observed when 10% ethanol was added to the water. Full removal at a low concentration can be achieved within the first few minutes of photoexcitation. The high-concentration solution shows a removal efficiency of 96.8% after 60 min. The degradation efficiency of U(vi)-ARZ3 complexes with a concentration of up to 100 mg L⁻¹ approaches almost 100% in a few minutes. The removal

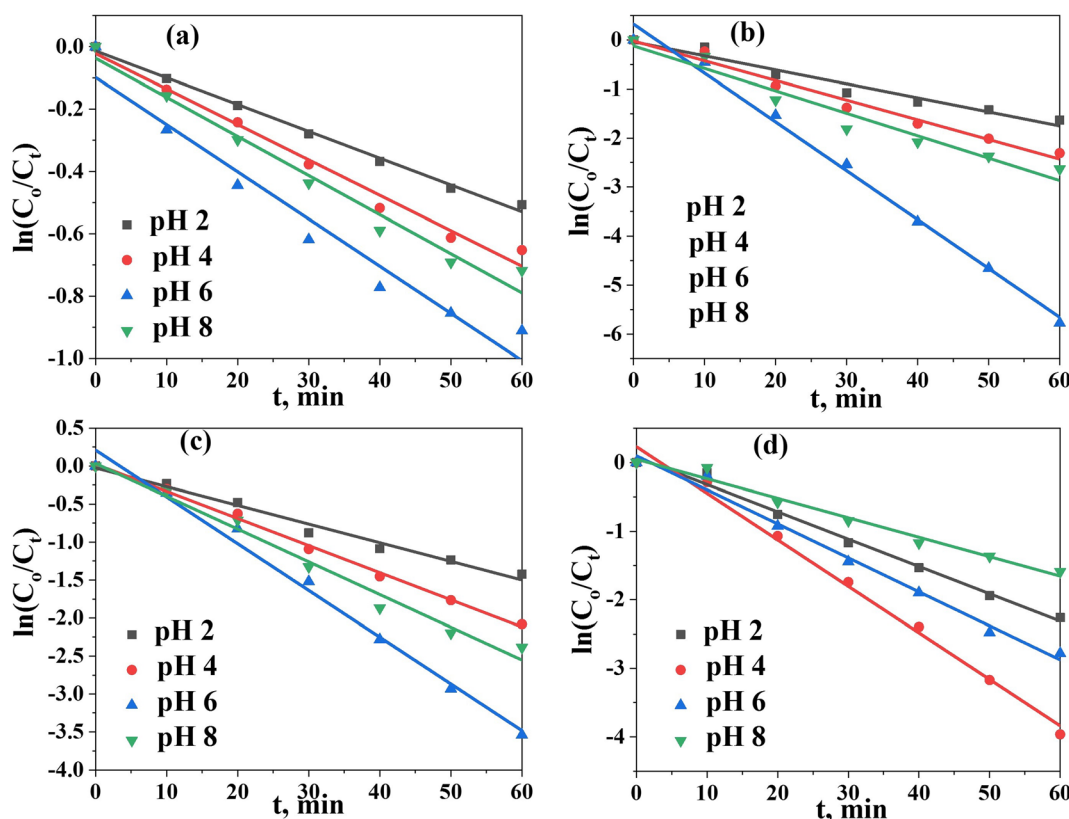


Fig. 6 Pseudo-first-order kinetic model. (a) U(vi) in aqueous solution, (b) U(vi) in aqueous solution (10% ethanol), (c) the U(vi)-ARZ3 complex, and (d) free ARZ3.



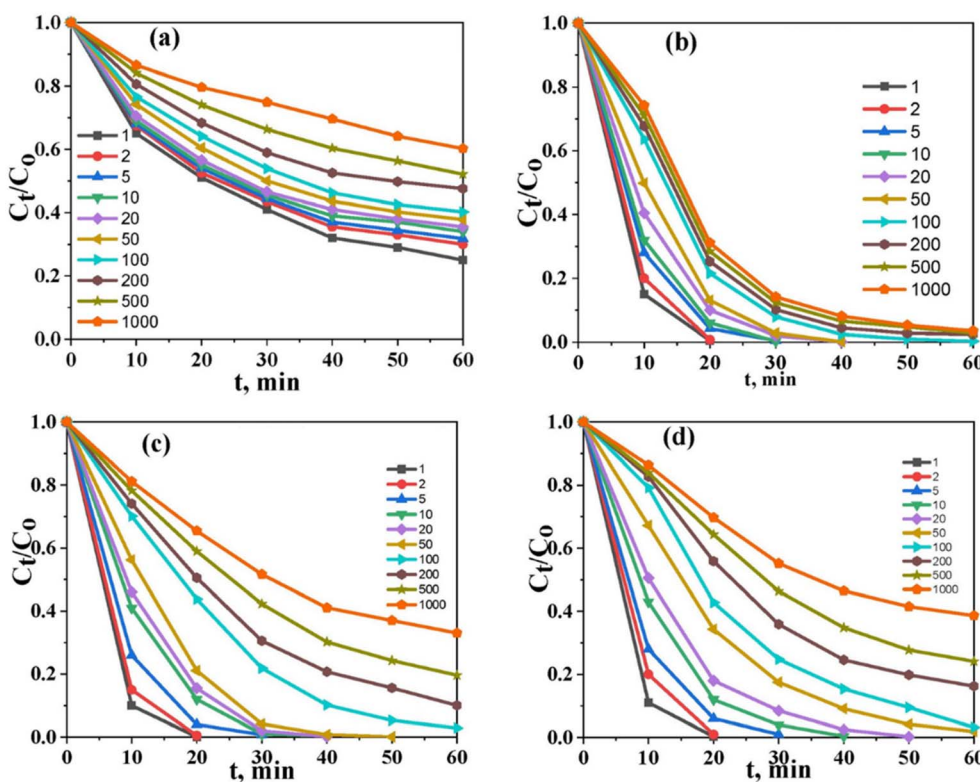


Fig. 7 The effect of the initial concentration under optimum conditions, (a) U(vi) in aqueous solution, (b) U(vi) in aqueous solution (10% ethanol), (c) the U(vi)-ARZ3 complex, and (d) free ARZ3.

efficiency decreases with an increase in the concentration, reaching 66.9% at a concentration of 1000 mg L^{-1} . Overall, the catalyst's efficiency is high at low concentrations and decreases with an increase in the concentration. We believe this is because of crowding the surface of the photocatalyst by new adsorbed ions and the need to remove the photodegraded product materials.

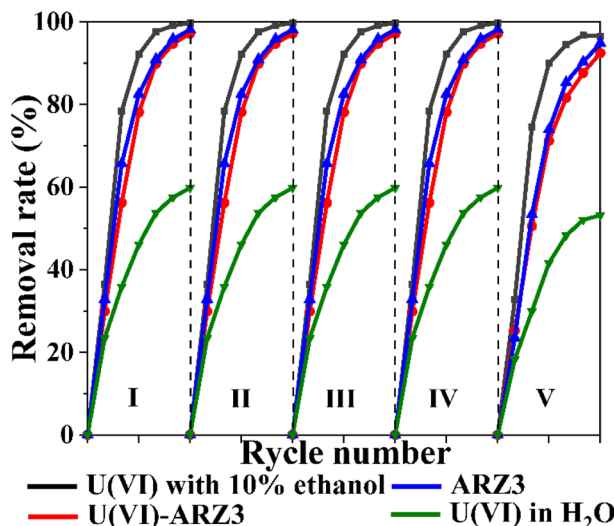


Fig. 8 Reusability studies of the Co@TiO₂-C catalyst [10 mL of 100 mg L⁻¹ concentration; 0.01 gm of catalyst; UV irradiation time, 60 min].

The reusability of catalysts is important from a practical point of view. Therefore, we have tested the Co@TiO₂-C catalyst for five successive cycling runs using U(vi)-ARZ3 solution (Fig. 8). Following each cycle, the Co@TiO₂-C was regenerated using 0.1 M HCl leaching, followed by water and ethanol washing. After the regeneration cycles, the degradation/reduction rate of uranium/ARZ3 remained almost consistent. The XRD spectra of Co@TiO₂-C before and after 5 runs show almost no changes, suggesting the stability of the catalyst (Fig. 1a). These results demonstrate the catalyst's stability and the possibility of using it for several removal cycles.

A comparison was made between the efficiency of the Co@TiO₂-C photocatalyst and some other catalysts that were used in the process of reducing dissolved U(vi) to insoluble U(IV) to remove nuclear pollutants from wastewater (Table. S2†).

After excitation, the photocatalytic products were recovered by centrifugation, dried in the air, and studied using SEM and EDX. Uranium and arsenic are detected by elemental mapping in addition to the primary components of Co@TiO₂-C (titanium, cobalt, carbon, and oxygen), as shown in Fig. S3–S5.† The XPS survey of the degraded product also confirmed the presence of arsenic and uranium in the precipitate after the photocatalytic process. Peaks for $\text{U } f_{5/2}$, $\text{U } f_{7/2}$, and $\text{As } 3d$ with a decrease in the intensity of $\text{Ti } 2p$, $\text{Co } 2p$, $\text{O } 1s$, and $\text{C } 1s$ (Fig. S6a and S7a†) can be observed. The high-resolution U 4f spectrum of the photocatalytic product in water/ethanol (Fig. 9a) shows two peaks at 393.54 eV and 384.95 eV, corresponding to $\text{U } 4f_{5/2}$ and $\text{U } 4f_{7/2}$ of U(vi),⁴² as well as two peaks at



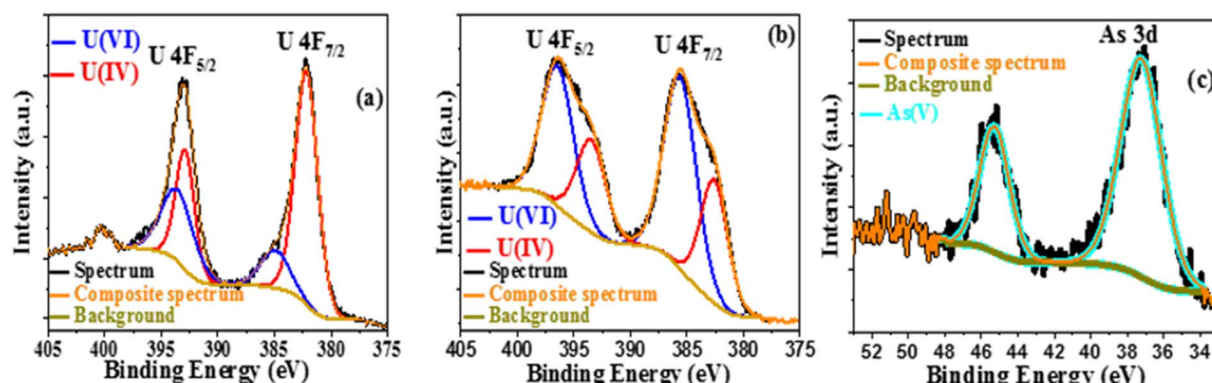


Fig. 9 (a) U 4f XPS spectra in 10% ethanol and (b and c) U 4f and As 3d XPS spectra of the U(vi)-AZR3 complex.

392.89 eV and 382.81 eV for U 4f_{5/2} and U 4f_{7/2} of U(IV).⁴² The same peaks can be observed for the product collected from the U(vi)-ARZ3 solution (Fig. 9b). Finally, Fig. 9c shows two peaks at 37.25 eV and 45.28 eV that can be ascribed to As 3d of As(v).⁴² The existence of hierarchical forms of arsenic oxide is visible in the SEM image, while uranium oxide appeared as short rod-like particles distributed across the surface of the Co@TiO₂-C catalyst (Fig. 10).⁸⁰

3.3. Photocatalytic mechanism

As discussed earlier, the crystal structure of anatase TiO₂ is distorted by Co and oxygen vacancies. Ti⁴⁺ is replaced by Co³⁺ and loaded onto graphite sheets, which in addition to the oxygen vacancies, produces a bandgap of 1.7 eV. Thus, upon irradiation, excitation occurs at the bandgap of Co/TiO₂, so the

photo-generated electrons (e⁻) move from the valence band (VB) to the conduction band (CB), leaving positive holes (h⁺) in their place on the VB, and then this e⁻ moves freely along the network of graphitic sheets. The graphitic sheets hinder electron and hole recombination,⁸¹ which facilitates charge separation and thus increases Co@TiO₂-C efficiency.⁸² On the surface of the graphitic sheets, the photo-generated electrons interact with O₂ to form 'O₂⁻', which can then react with H₂O to form 'OH/H₂O₂'.⁸³ In addition, h⁺ produced in the VB may react with H₂O in the medium to form 'OH'. This free radical 'O₂⁻'/ 'OH and photo-generated e⁻/h⁺ can oxidize ARZ3 and reduce U(vi) to U(iv). The effects of free radicals on photocatalytic U(vi) reduction were further verified *via* radical trapping experiments, where ethanol and ARZ3 were added as scavengers for free radical 'OH and photo-holes h⁺. Comparing the results in

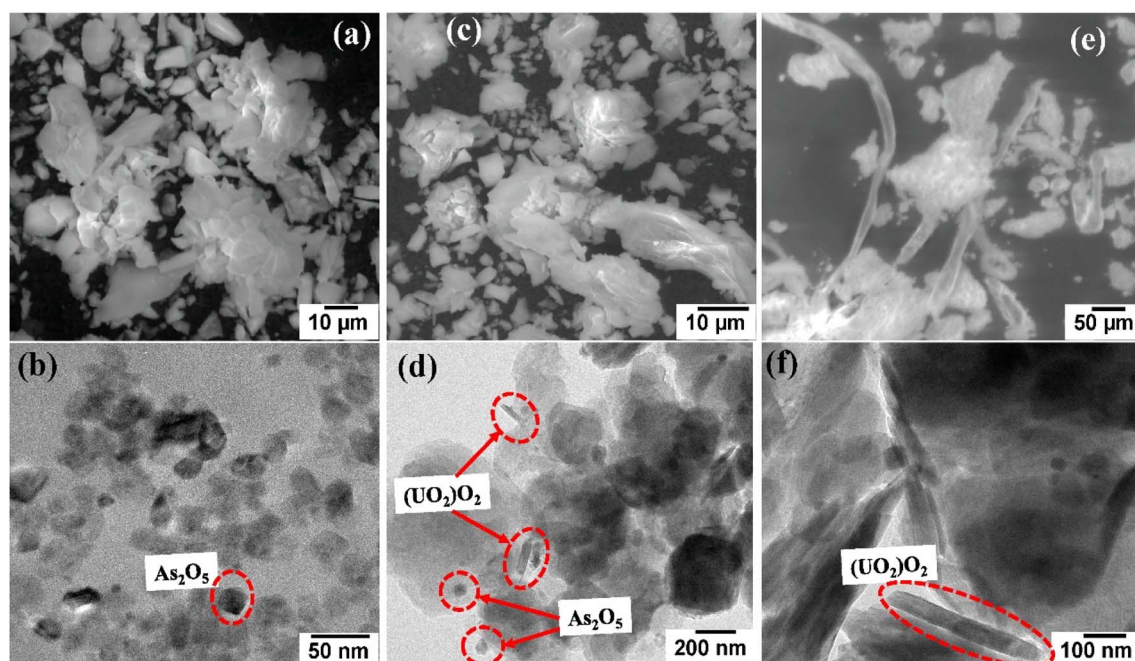


Fig. 10 (a and b) SEM and TEM images of the ARZ3 degraded product. (c and d) SEM and TEM images of uranium and the arsenazo III degraded product. (e and f) SEM and TEM images of the uranium degraded product in ethanol media.



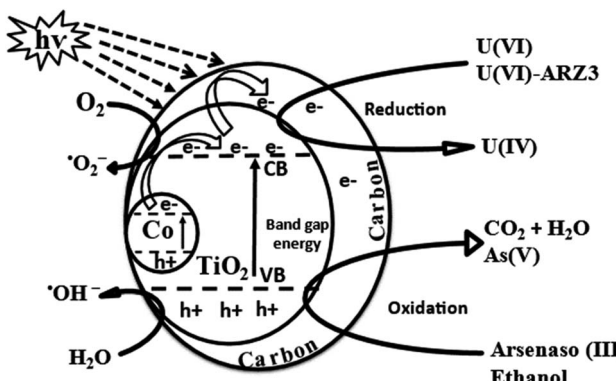
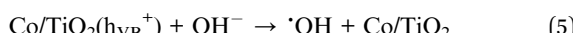
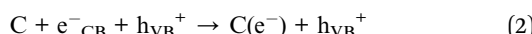
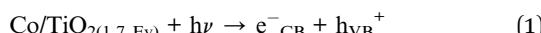
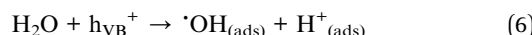


Fig. 11 Photocatalytic reduction of the U(vi)-ARZ3 mechanism using C@TiO₂-C under UV light.

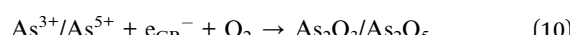
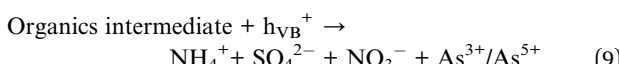
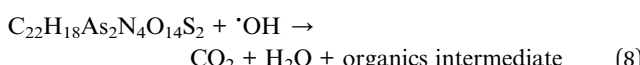
Fig. 4c indicates that in the absence of ethanol and ARZ3, U(vi) reduction has been reduced, while the presence of ethanol and U(vi)-ARZ3 promotes the removal of U(vi). These results demonstrated that ·O₂-radicals acted as the key reductive species for U(vi) reduction, while ·OH radicals had negative effects on the reduction of U(vi).⁸⁴ As a result, arsenic oxide and uranium oxides precipitate out of the solution, in agreement with the EDX results. This reaction mechanism can be described by using eqn (1) through (15) and is schematically summarised in Fig. 11.



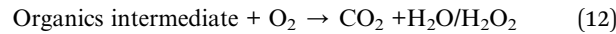
H₂O



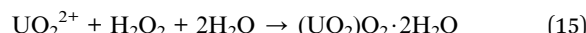
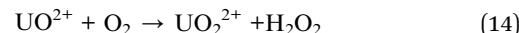
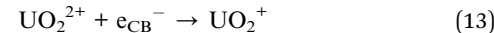
ARZ3



Ethanol



U(vi)



4. Conclusion

A carefully designed photocatalyst (Co@TiO₂-C) was used to remove radioactive wastes under UV light. The photocatalytic performance of cobalt-doped titanium oxide was improved by inducing oxygen vacancies in the lattice and controlling the structure into 2D nanosheets. Hybridizing the oxide nanostructures with carbon during the hydrothermal process helped to prevent the recombination of the photo-generated holes and electrons and, in addition, to improve the catalytic efficiency, allowing the degradation of uranium complexes commonly associated with radioactive waste. Also, benefiting from the high surface area of the Co@TiO₂-C catalyst, the photo-degradation process was fast, with the ability to remove most of the uranium complex within 20–30 minutes of the process. The reaction mechanism was also investigated, and the spectroscopic analysis confirmed the reduction of soluble U(vi) to insoluble U(iv). It was also found that adding ethanol to the aqueous solutions improves the photocatalytic process by working as a scavenger to the photo-generated hole and minimizing the formation of hydrogen gas bubbles on the surface of the catalysts. We believe that the photocatalysts used in this work, coupled with the simple and affordable synthesis method, could provide new strategies and insights for the photo-reduction of toxic ions from polluted water.

Conflicts of interest

There are no conflicts to declare.

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