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Competitive photooxidation of small colorless organics controlled by oxygen vacancies under visible light†

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Visible-light photooxidation sensitized by surface attachment of small colorless organics on semiconductor photocatalysts has emerged as an economical method for photocatalytic synthesis or degradation. In particular, heteroatom (X = N and Cl)-containing substrates could undergo either C–N coupling or dechlorination degradation *via* sensitizing TiO₂, but the mechanism in conducting the competitive visible-light sensitized photooxidations is still vague. Herein, the visible-light photooxidation of colorless 4-chlorobenzene-1,2-diamine (*o*-CAN) on TiO₂ was revealed, contributing to selective C–N coupling rather than dechlorination. Oxygen vacancies (OVs) were *in situ* generated on the TiO₂ surface, which could be dominant in weakening the Cl–Ti adsorption of *o*-CAN and regulating the activation of O₂ for selective C–N coupling. The C–N coupling product, functionalized as the sensitizer, further promoted the visible-light photooxidation upon N–Ti and Cl–Ti coordination. This process was then confirmed by on-line mass spectrometric analysis, and the intermediates as well as their kinetics were determined. Thereby, theoretical calculations were employed to verify the roles of OVs in competitive photooxidation and lowering the energy barriers as well. Based on the comprehensive characterizations of both the catalysts and intermediates, this work has provided insights into competitive photooxidations under visible light.

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Introduction

Visible light catalysis stimulated by surface attachment of small colorless organics on semiconductor photocatalysts has become an economical and environmental route for chemical transformations.^{1–4} The easy adsorption of heteroatom (X = O, S, N or Cl)-containing organics on TiO₂ greatly promotes the visible-light photooxidation because the ensuing hybrid interfaces allow electron injection from electron rich heteroatoms to the TiO₂ conduction band.^{5–7} With O₂ as the terminal oxidant rather than undesirable sacrificial electron donors (like TEMPO),^{8,9} different oxidation processes would be induced *via* regenerating the surface-bound sensitizer molecules.¹⁰ For example, colorless benzylic amines can be oxidized into the

corresponding imines *via* the formation of a surface complex.¹¹ Meanwhile, the degradation of chlorinated pollutants can also be facilitated by themselves or stable photocatalytic sensitizers.¹² Nevertheless, most reports normally focus on developing catalysts or sensitizers for selective photooxidation, and the competitive mechanism between C–N coupling and dechlorination degradation on sensitized catalysts is still vague. Consequently, clear insights into the photooxidations are required, including competitive adsorption of substrates, activation of oxidants (like O₂), interactions of the sensitizer with the catalysts and the process of molecular transformation.

Notably, native oxygen vacancies (OV) on the surface of metal oxides is an important defect to mediate catalytic reactions *via* ameliorating photocatalysts.^{13–15} For example, OVs could be reactive sites to increase charge separation/migration and regulate the adsorption of substrates. In addition, OVs also facilitate the decrease of the reaction barrier to promote substrate activation,¹⁶ intermediate formation¹⁷ or byproduct suppression.¹⁵ In particular, OVs are conducive to the adsorption and activation of O₂, which could either be reduced to superoxide (O₂^{•−}) or incorporated into the product for oxo-functionalization.¹⁸ Significantly, the generated O₂^{•−} could induce amine oxidation through two different reaction pathways: (1) participating in the oxidative dehydrogenation to afford hydrogen peroxide (H₂O₂) during C–N coupling.^{19,20} (2)

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Forming hydroxyl radicals ($\cdot\text{OH}$) to initiate degradations for obtaining dechlorination products or even smaller molecules (CO_2 and H_2O).^{21,22} Consequently, strategies for creating more OVs on semiconductors have boomed, including doping, reduction and calcination. However, a report revealing the roles of OVs in visible-light sensitized competitive photooxidations is still lacking. Therefore, detailed studies of OV-controllable competitive photooxidation are required, which would be associated to the regulation of molecular adsorption and O_2 activation on sensitized semiconductor photocatalysts.

Herein, the visible-light photooxidation of colorless 4-chlorobenzene-1,2-diamine (*o*-CAN) on a TiO_2 surface was comprehensively examined by both off-line and on-line examinations. Interestingly, even in the presence of $-\text{Cl}$ (a potential degrading groups),¹² *o*-CAN still exhibited selective C–N coupling to generate 7-chlorophenazine-2,3-diamine (7-Cl-Phz) under mild conditions,²³ without necessary substrate pre-functionalizations.^{24–27} With the aid of *in situ* generated OVs on TiO_2 , the produced 7-Cl-Phz acted as the TiO_2 photosensitizer to further facilitate C–N coupling rather than degradation. The selective C–N coupling was attributed to OVs *via* promoting interfacial electron transfer, regulating *o*-CAN adsorption and suppressing $\cdot\text{OH}$ generation. Furthermore, online mass spectrometry analysis was employed for structural identification and dynamic monitoring of intermediates, being conducive to revealing the roles of OVs. As demonstrated by experimental and theoretical examinations, selective C–N coupling was obtained by manipulating substrate adsorption, O_2 activation and energy barriers by OVs. This work would not only promote the development of controllable and selective visible-light photooxidation, but also inspire a deeper understanding of nanocatalysis.

Results and discussion

Evaluation of *o*-CAN visible-light photooxidation

The photooxidation of N, Cl-containing *o*-CAN catalyzed by TiO_2 was selected as the model for examinations. In the reaction, commercial P25- TiO_2 was added into the *o*-CAN solution, following by the visible-light irradiation under mild conditions. Under visible-light irradiation, the colorless *o*-CAN (with $-\text{NH}_2$ and $-\text{Cl}$ groups) could be oxidized into 7-Cl-Phz *via* C–N coupling, or degraded into radical or 3,4-diaminophenol (DAP) in the presence of $\cdot\text{OH}$ (Fig. 1A). The reaction system was coupled with the multiphase flow of extractive electrospray ionization mass spectrometry (MF-EESI) for reaction monitoring.²⁸ As shown in Fig. 1B, the reaction solution was suspended in a double-decker quartz beaker equipped with a temperature controller. The reaction system (RS) was then coupled to the ionization system (IS) for on-line reaction monitoring. According to our previous studies,^{23,29} analytes would be extracted by the cooperation of inserted capillary and high-speed N_2 . Simultaneously, the concentric-spray of methanol and a high DC voltage can facilitate the ionization upon decreasing the interference of TiO_2 .

As demonstrated (Fig. 1C(i)), after irradiating for 25 min, the main ion peak was ascribed to the C–N coupling product of [7-

Cl-Phz + H]⁺ at m/z 245, which was further confirmed by collision-induced dissociation (CID) (Fig. S1†) and ^1H NMR (Fig. S2†) analyses. On the contrary, no significant ions associated with the degradation were observed, and nor was the dechlorinated ion of *o*-PD⁺ at m/z 108 (the structure was identified by MS/MS analysis in Fig. S3†) or the $\cdot\text{OH}$ -related degradation product of DAP in both the positive (m/z 125, the dashed) and negative mode (m/z 123, Fig. S4†).³⁰ It should be noted that to avoid decomposition of the active PD⁺ before detection, the distance from the spray tip to the MS inlet was set at 0.5 cm (Fig. S5†). Besides, a relatively low abundance of reactant ion ([*o*-CAN + H]⁺ at m/z 143) was exhibited, indicating the efficient and selective C–N coupling reaction. Consequently, upon photosensitization of TiO_2 under visible-light irradiation, the colorless *o*-CAN preferred to undergo selective C–N coupling rather than degradation.

Furthermore, the dynamic changes of the dechlorination product of *o*-PD⁺ and C–N coupling product of 7-Cl-Phz were on-line monitored by MF-EESI. The online extracted ion chromatograms (EICs) of *o*-PD⁺ (m/z 108) and [7-Cl-Phz + H]⁺ (m/z 245) were collected for comparison. As demonstrated (Fig. 1C(ii)), the photooxidation could be divided into three stages. Before approximately 1.5 min (S_1), quite low ion signals were recorded. Although a slightly higher signal of [7-Cl-Phz + H]⁺ than *o*-PD⁺ was recorded, either the degradation or C–N coupling was non-selective. Thereafter (S_2), [7-Cl-Phz + H]⁺ dramatically increased along with the gradual decrease of *o*-PD⁺, followed by the kinetic equilibrium of both signals at ~ 20 min. After 20 min (S_3), [7-Cl-Phz + H]⁺ decreased slightly along with the generation of dechlorinated Phz⁺ (m/z 210, the inset of Fig. 1C(i) and S6†), while no significant change of *o*-PD⁺ was observed. These results indicate two possible facts: (1) the easier absorption of the N,Cl-containing sensitizer of 7-Cl-Phz on TiO_2 enhanced photoinduced electron injection, while weaker conjugation of *o*-CAN or *o*-PD⁺ with TiO_2 disfavored the photosensitization and further degradation. (2) Degradation of *o*-CAN was further blocked by the accumulation of the 7-Cl-Phz sensitizer/product on TiO_2 .

The photooxidation of *o*-CAN catalyzed by TiO_2 was further evaluated based on the significant UV-vis absorption of 7-Cl-Phz in the visible light region (~ 445 nm, Fig. S7-A†). Compared to the absorption of the present *o*-CAN photooxidation (Fig. S7-B,† entry 1), a similarly high response was obtained by extra addition of 7-Cl-Phz to support photosensitization (Fig. S7-B,† entry 2). Besides, a series of control experiments were carried out to examine the efficient and selective photocatalytic C–N coupling. As demonstrated, the reaction was hardly carried out in the absence of either TiO_2 or light irradiation (Fig. S7-B,† entries 3 and 4). Notably, a low yield of 7-Cl-Phz was produced without O_2 (Fig. S7-B,† entry 5), indicating that O_2 could be the terminal oxidant for *o*-CAN photooxidation.

Furthermore, to examine the intrinsic effect of the substrate for selectivity, a scope investigation with diverse structures has been carried out. As demonstrated, without a $-\text{Cl}$ group, only the C–N coupling product was observed when using *o*-phenylenediamine or 4-methyl-*o*-phenylenediamine as the substrate (Fig. S8a and b†). Without $-\text{NH}_2$ groups, the chlorobenzene was



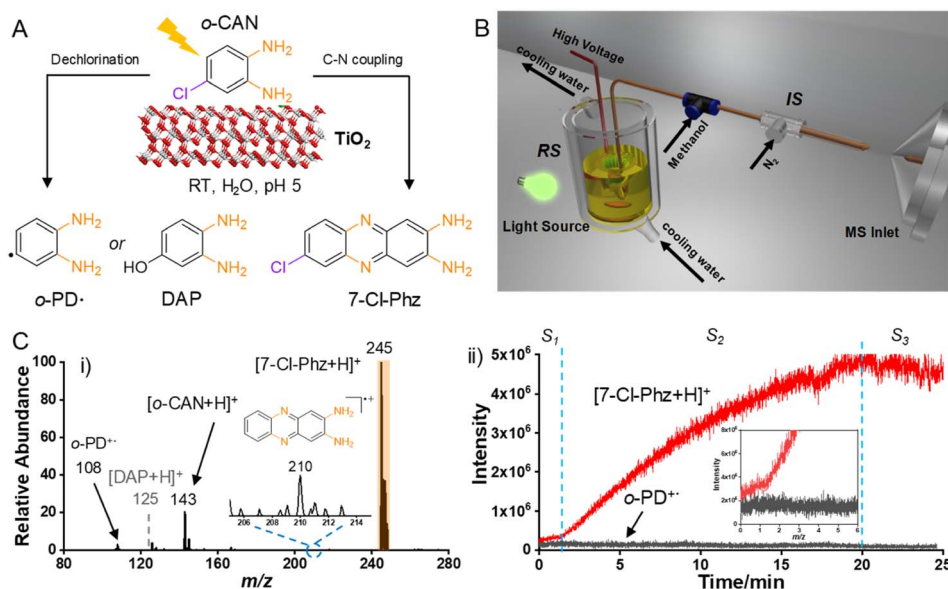


Fig. 1 Schematic diagram and reaction evaluation via online mass spectrometry. (A) Schematic diagram of photocatalytic C–N coupling and degradation on TiO₂. (B) Schematic diagram of the online reaction monitoring system by coupling the reaction system with an ionization system.²⁸ (C) (i) Mass spectra of the reaction system after visible-light irradiation for 25 min in air. Reaction conditions: α-CAN (0.5 mM), TiO₂ (1 mg mL⁻¹), H₂O (20 mL, pH = 5), and a xenon lamp (simulated sunlight, AM 1.5). (ii) EICs of the C–N coupling product of [7-Cl-Phz + H]⁺ and the degradation product of o-PD⁺.

degraded into ketone or aldehyde (Fig. S8c†), indicating the feasibility of dechlorination degradation. Besides, both the C–N coupling product ions and corresponding degradation product ions were detected for other substrates with –Cl at different positions and –Br substitution, demonstrating the occurrence of competitive reactions (Fig. S8d and e†). When increasing the number of –Cl, the dechlorination degradation was promoted, but a small amount of C–N product ion (*m/z* 313) was still observed (Fig. S8f†), especially considering the steric hindrance of the –Cl groups. This demonstrated the negative role of –Cl in the C–N coupling reaction, which could be promoted by *in situ* generated OVs.

Consequently, the generated 7-Cl-Phz can induce dye/self-sensitization of TiO₂ to facilitate the competitive photooxidation of α-CAN, which turned out to further promote the C–N coupling. However, this is different from the reports, which demonstrated that the sensitization-based visible-light catalysis of TiO₂ is non-selective and could also facilitate degradation.^{31,32} Consequently, more detailed mechanism studies are required to reveal the competitive photooxidation mechanism in conducting the selective C–N coupling, rather than degradation.

Surface engineering of TiO₂

Chemical changes of TiO₂ after the photooxidation were characterized. Demonstrated by high resolution transmission electron microscopy (HR-TEM) (Fig. S9†), the particle size of TiO₂ increased after the reaction, along with a decreased Brunauer–Emmett–Teller (BET) surface area (*S*_{BET}). The visible light absorptions of TiO₂ before and after the reaction were further examined by diffuse reflectance UV-vis (DR-UV-vis) spectra analysis. As demonstrated (Fig. 2A), a broader and extended

absorption of TiO₂ (from 400 to 700 nm) was recorded after the reaction. The above results indicated the adsorption of 7-Cl-Phz on TiO₂ (Fig. S7-A†), which could act as the photosensitizer to broaden the spectral responses of semiconductors with a wide-band gap. This was also consistent with the reported easy absorption of heteroatom-containing dyes (or substrates) onto the TiO₂ surface.³³

The strong interaction between TiO₂ and 7-Cl-Phz was further confirmed by X-ray diffraction (XRD) characterizations. As illustrated in Fig. 2B, the XRD peaks were red-shifted after the photooxidation, which was attributed to a lattice expansion caused by surface modification of heteroatoms (N or Cl) into the unit cell.³⁴ To further examine the surface chemical composition of TiO₂, X-ray photoelectron spectroscopy (XPS) characterizations were employed and the corresponding evident peaks were deconvoluted in the HR-XPS spectra. As demonstrated, 7-Cl-Phz exhibited the characteristic peaks of C=N–C and C–NH₂ at binding energies of 398.94 and 399.91 eV in the N 1s spectrum (Fig. 2C(a)-i).³⁵ Meanwhile, after the reaction (Fig. 2C(a)-ii), the two peaks shifted to a lower binding energy (398.84 and 399.51 eV), along with the recording of the N–Ti–O peaks (397.99 and 400.34 eV).³⁶ Similarly, the original Cl 2p_{3/2} and Cl 2p_{1/2} peaks of 7-Cl-Phz (at 200.07 and 201.55 eV in the XPS Cl 2p spectra, Fig. 2C(b)-i) shifted to higher binding energies (at 200.45 and 201.98 eV) along with the formation of Cl–Ti and Ti–Cl–Ti bonds (197.58 and 199.80 eV, Fig. 2C(b)-ii).^{37,38} Besides, the characteristic peaks of reformed Ti–Cl and Ti–N–O bonds (at 457.66 and 463.17 eV) were recorded after the reaction (Fig. S10†),^{39,40} which was in agreement with the N 1s and Cl 2p data (Fig. 2C). Consequently, the adsorption of 7-Cl-Phz on the TiO₂ surface was further confirmed upon the direct bonding of both hetero atoms (Cl and N) to Ti.



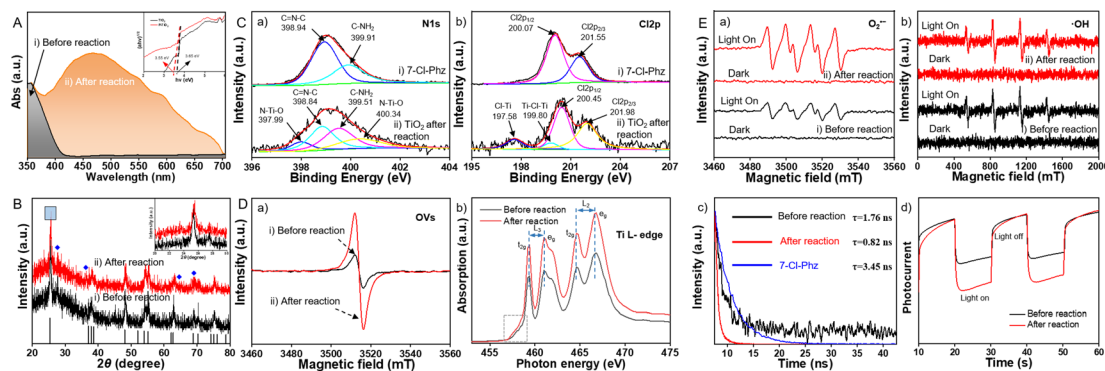


Fig. 2 Chemical characterizations of TiO₂ before and after the photooxidation. (A) DR-UV-vis spectra. (B) XRD spectra. (C) Core level HR-XPS spectra in the N 1s region (a) and Cl 2p region (b). (D) EPR spectra of OVs (a) and XAS spectra (b). (E) EPR spectra of O₂^{•-} in the dark and under photoirradiation (a) and [•]OH in the dark and under photoirradiation (b); (c) TRPD spectra of 7-Cl-Phz and TiO₂ before and after the photooxidation; (d) transient photocurrent responses. Conditions: +0.5 V versus Ag/AgCl at pH 5.0. Irradiation: 300 W Xe lamp of simulated solar light (100 mW cm⁻²), under chopped illumination with a fixed time interval.

Subsequently, the surface defects of oxygen vacancies (OVs) were examined by electron spin resonance (ESR) and soft X-ray absorption fine structure (sXAFS) analysis. After the photocatalytic oxidation (Fig. 2D(a)), the EPR signals of the OVs on TiO₂ increased significantly, indicating the *in situ* formation of OVs along with the absorption of the 7-Cl-Phz sensitizer. The formation of OVs could be facilitated by the abundant surface -OH, acting as Brønsted base sites for reductive deprotonation (oxidation) of aniline.^{41,42} Furthermore, the local electronic configuration of TiO₂ was further studied by Ti L-edge. Upon photooxidation (Fig. 2D(b)), the onset energy position of the Ti L-edge of TiO₂ shifted to a lower energy, suggesting a lower Ti valence state empirically.⁴³ As demonstrated, along with the adsorption of the 7-Cl-Phz sensitizer, surface defects of OVs were *in situ* generated during the photooxidation of *o*-CAN. In fact, the generation of surface OVs was crucial for the competitive photooxidation, which was confirmed by the increased C-N coupling efficiency with increasing the concentration of OVs on the TiO₂ surface (Fig. S11[†]).

To further examine the role of OVs, characterizations including ESR, time-resolved photoluminescence decay (TRPD) and electrochemical impedance spectroscopy (EIS) were employed. As demonstrated (Fig. 2E(a)), O₂^{•-} was significantly generated on TiO₂ after the reaction, induced by the efficient reduction of O₂ upon capturing photoelectrons. Meanwhile, no significant increase of [•]OH was observed (Fig. 2E(b)), indicating that no further degradation would be employed after the generation of *o*-PD[•] via dichlorination of *o*-CAN (as illustrated in Fig. 1A). In addition to O₂^{•-} and [•]OH, reactive oxygen species of O₂¹ were also detected, while no signal was observed on TiO₂ even after the reaction (Fig. S12[†]). Therefore, the *in situ* generated OVs could modulate O₂ activation and electron transfer to promote the selective C-N coupling.

Besides, the interfacial charge transfer of TiO₂ before and after the reaction was evaluated. Firstly, the TRPD spectra (Fig. 2E(c)) were collected to fit the decay traces, which demonstrated a distinctly faster decay of TiO₂ after the reaction ($\tau_{\text{average}} = 0.82$ ns) than that before the reaction ($\tau_{\text{average}} = 1.76$

ns).^{44,45} Consequently, the rapid charge transfer between the absorbed 7-Cl-Phz and the conduction band (CB) of TiO₂ was confirmed, which facilitated the subsequent photooxidation under visible light. Besides, the excited-state lifetime of 7-Cl-Phz was 3.45 ns, much longer than that of TiO₂ before the reaction (1.76 ns). This indicated that there would be sufficient time for electron transfer from the excited state of 7-Cl-Phz to TiO₂ before relaxation. Furthermore, photocurrent and electrochemical impedance were also evaluated. An increased photocurrent density of TiO₂ was observed after the reaction (Fig. 2E(d)), indicating a more efficient photogenerated electron-hole separation on the interface of [7-Cl-Phz]-TiO₂. The ensuing active holes on the hetero atoms would promote the subsequent photooxidation of *o*-CAN. Correspondingly, a decreased R_{rec} value of TiO₂ after the reaction (from 2091 Ω to 1976 Ω) was also recorded upon fitting the Nyquist plot (Fig. S13[†]). Therefore, the adsorption of 7-Cl-Phz on TiO₂ could alter the electronic structure and defect states to enhance charge transfer and separation. This property would be highly beneficial for efficient and selective photooxidation on semiconductor catalysts with wide-band gaps.

Theoretical examinations of OV-controlled competitive photooxidation

To explore the roles of OVs in the competition, theoretical calculation was performed by density functional theory (DFT) using the Vienna *Ab Initio* Simulation Package (VASP).^{46,47} Firstly, the adsorptions of the reactant (*o*-CAN) on TiO₂ through both Ti-N and Ti-Cl modes without OVs, as well as that on TiO₂ with abundant OVs (TiO₂-OVs) were evaluated for comparison. Without OVs, *o*-CAN was stably adsorbed on TiO₂ in both the Ti-N and Ti-Cl modes (Fig. 3A(i)). Meanwhile, in the presence of OVs, a higher adsorption energy of Ti-Cl (0.4 eV) than Ti-N (-0.13 eV) was exhibited (Fig. 3A(ii)). This result indicated that OVs could weaken the direct interaction of Ti-Cl to hinder the dichlorination. The differential charge density of *o*-CAN and ^{*}*o*-CAN on TiO₂ in Ti-Cl modes further verified the limitation of Ti-Cl adsorption (Fig. S14[†]). Compared to the *o*-CAN molecule,



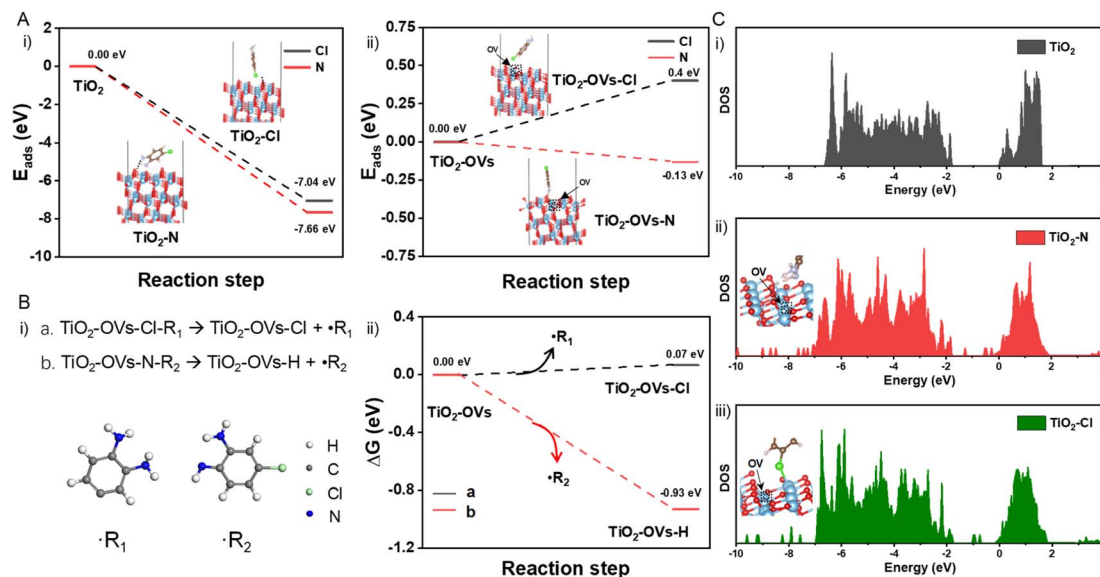


Fig. 3 Theoretical calculation of competitive photooxidation for examining selective C–N coupling. (A) Adsorption energies of *o*-CAN on TiO₂ (i) and TiO₂-OVs (ii). (B) (i) Illustration of the dechlorination and dehydrogenation to generate ·R₁ and ·R₂, respectively. (ii) Free energy changes of dechlorination and dehydrogenation on TiO₂-OVs. (C) DOS of the metal d orbital before (i) and after adsorption via Ti–N (ii) and Ti–Cl (iii).

more electron depletion (cyan) of the N atom of *o*-CAN in the Ti–Cl mode was observed. This indicated the decreased electron density of the N atom after adsorption in Ti–Cl mode, which could hinder its further oxidation for selective C–N coupling.

Besides, the energy change of the initial dechlorination (for oxidative degradation) and dehydrogenation (for C–N coupling) (Fig. 3B(i)) was further evaluated. As shown in Fig. 3B(ii), the energy change of degradation (to release ·R₁) is much higher (0.07 eV) than that of the C–N coupling (to release ·R₂) (–0.93 eV). Meanwhile, higher energy barriers for both the degradation (2.3 eV) and C–N coupling (2.0 eV) are produced when catalyzed by TiO₂ without OVs (Fig. S15[†]), further confirming the effect of OVs on selectivity. Consequently, promoted by the *in situ* generated OVs, the oxidative dehydrogenation of *o*-CAN (for C–N coupling) is more favorable than the dechlorination (for oxidative degradation). Subsequently, to investigate the photo-induced charge transfer (PICT) between the 7-Cl-Phz sensitizer and TiO₂, DFT was employed to calculate the electronic density of states (DOS) of TiO₂ before and after the photooxidation. As shown, direct bonding of 7-Cl-Phz *via* both Ti–N (Fig. 3C(ii)) and Ti–Cl (Fig. 3C(iii)) rendered smaller band gaps than that of the crystal counterpart (Fig. 3C(i)). Although the DFT results of semiconductor structures are qualitative assessments, it is still sufficient to predict the enhancement of PICT by the bonding between 7-Cl-Phz and TiO₂.

Reaction monitoring and kinetic studies

To explore the molecular changes and the generation of intermediates, the photooxidation was monitored by *in situ* Raman spectroscopy.^{48–51} As shown in Fig. 4A, upon 1 min of irradiation, only typical Raman bands (B_{1g(1)}, A_{1g} + B_{1g(2)}, and E_{g(2)}) of TiO₂ were recorded. This indicated the weak bonding between the substrate (*o*-CAN) and TiO₂ with lower visible-light

absorption and inefficient electron transfer. After 3 min irradiation, the peaks of ν_{CCl} (1026 cm^{–1}), ν_{NH} (1229 cm^{–1}) and ν_{CN} (1580 cm^{–1}) increased upon the accumulation of the 7-Cl-Phz product. Thereafter, a new peak at 1518 cm^{–1} appeared and increased gradually, which was attributed to the formation of quinonoid intermediates. In addition, a shoulder peak of C~N⁺ (~ denotes a bond intermediate between a single and double bond) at 1330 cm^{–1} was also observed, attributed to the stretching vibration of polaronic structures. Significantly, half-oxidized phenazine-like units were also demonstrated by the strong peak at 1402 cm^{–1}, which could be assigned to the free radical precursor of 7-Cl-Phz.^{52,53} Consequently, the quinonoid intermediates, C~N⁺ intermediates and half-oxidized phenazine-like radical precursors were preliminarily confirmed to be involved in the photooxidation.

To directly obtain the molecular information during the selective C–N coupling, intermediate structures, dynamics of reactants, intermediates and products were determined by MS detections. As indicated by the MS spectra at the interval times (Fig. 4B(i)), only the reactant ion of *o*-CAN ([1+H]⁺) at *m/z* 143 was observed without light irradiation. Upon visible light irradiation (Fig. 4B(ii)), the product ion of 7-Cl-Phz ([7+H]⁺) at *m/z* 245 came to be observed, which would act as the antenna molecule for visible-light absorption. In addition, the multiple ions attributed to the protonated dimers of *o*-CAN were observed (Fig. 4B(iii–v)), including the ions at *m/z* 283 ([3+H]⁺, measured *m/z*: 283.0517, theoretical *m/z* 283.0512, error +0.6 ppm), *m/z* 281 ([4 + H]⁺, measured *m/z*: 281.0363, theoretical *m/z* 281.0355, error +0.79 ppm), *m/z* 249 ([6+H]⁺, measured *m/z*: 249.0910, theoretical *m/z* 249.0902, error +1.7 ppm), and *m/z* 247 ([5+H]⁺, measured *m/z*: 247.0757, theoretical *m/z* 247.0745, error +2.6 ppm). The corresponding structures are shown in the insets of Fig. 4B, which were confirmed *via* CID experiments



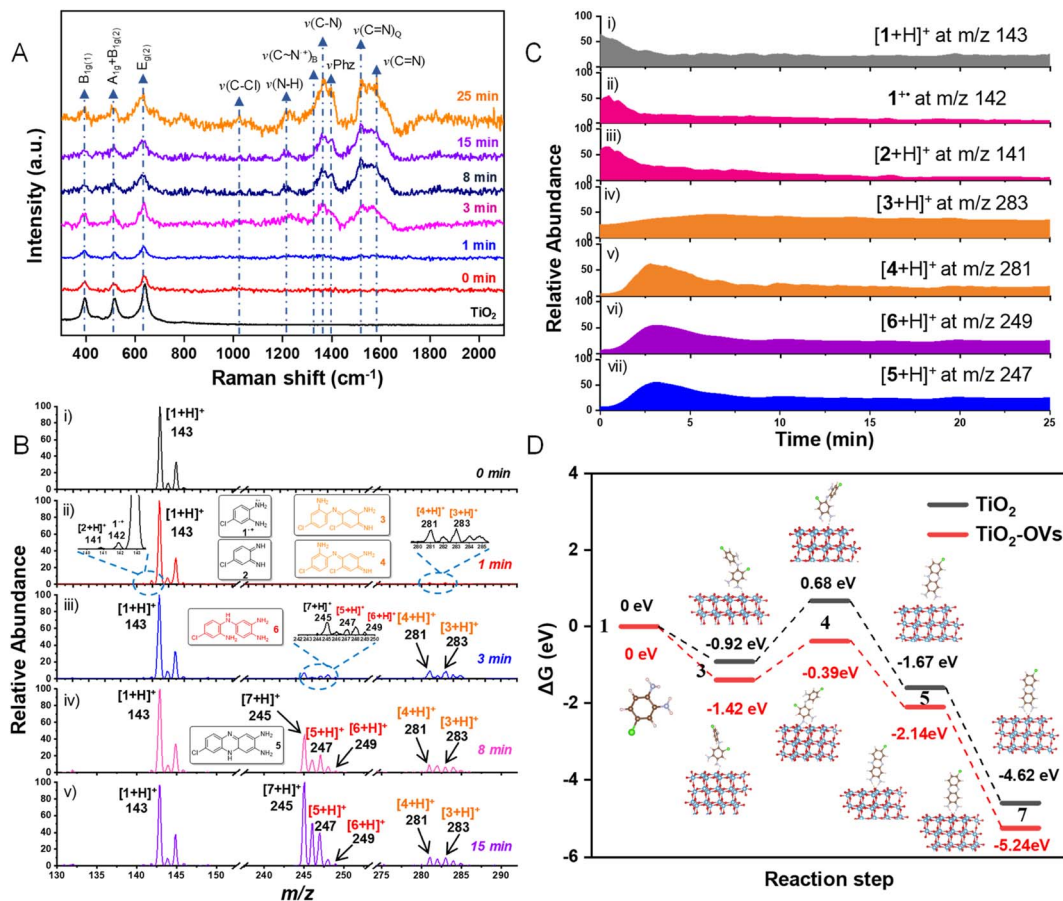


Fig. 4 Reaction monitoring. (A) *In situ* Raman spectroscopic monitoring of photooxidation at different times. (B) Mass spectra of the photooxidation reaction at different times *via* on-line MS monitoring. (C) EICs of [1+H]⁺ at *m/z* 143 (i), intermediate ions of 1⁺ (*m/z* 142) (ii), [2+H]⁺ (*m/z* 141) (iii), [3+H]⁺ (*m/z* 283) (iv), [4+H]⁺ (*m/z* 281) (v), [6+H]⁺ (*m/z* 249) (vi), and [5+H]⁺ (*m/z* 247) (vii). (D) Gibbs free energy diagrams for 1 → 3 → 4 → 5 → 7 on TiO₂ and TiO₂-OVs, respectively.

(Fig. S16 and S17[†]). The generation of these intermediates was further confirmed by separating the corresponding ion peaks from these isobars (Fig. S18[†]). Besides, the ions of C~N⁺ and the quinonoid intermediates at *m/z* 142 and 141 were also recorded upon light irradiation, in accordance with the Raman data in Fig. 4A. The corresponding structures were further confirmed by CID experiments (Fig. S19[†]). Significantly, the recorded ions exhibited different dynamic changes with time-dependent ratios of the relative abundance, which required further dynamic examinations by on-line monitoring.

Thereafter, the kinetics of the dye/self-sensitized photooxidation on TiO₂ were online monitored by MF-EESI MS and the extracted ion chromatograms (EICs) of different ions were collected. As shown in Fig. 4C(i), the reactant ion of [1+H]⁺ at *m/z* 143 decreased along with the reaction taking place. Meanwhile, the ions of 1⁺ (*m/z* 142) and [2+H]⁺ (*m/z* 141) increased immediately within 0.4 min (Fig. 4C(ii and iii)), which were generated from the rapid oxidation of 1 to form 1⁺ and then to 4-chlorocyclohexa-3,5-diene-1,2-diimine (2). Subsequently, the ions of 1⁺ and [2+H]⁺ gradually decreased, which indicated that both ions could be important intermediates of the photooxidation. Interestingly, along with the relatively rapid decrease of

[2+H]⁺ (Fig. 4C(iii)), [3+H]⁺ (*m/z* 283) increased gradually until 6.5 min (Fig. 4C(iv)). Simultaneously, [4+H]⁺ (*m/z* 281) increased dramatically and was rapidly consumed after 2.8 min (Fig. 4C(v)). Both ions of [3+H]⁺ and [4+H]⁺ could be assigned as the intermediates converted from 2. Simultaneously, the ion of [6+H]⁺ (*m/z* 249) also exhibited a gradual increase and decrease after ~3 min, indicating another potential pathway with 6 as the intermediate (Fig. 4C(vi)). Along with the generation of [4+H]⁺ and [6+H]⁺, [5+H]⁺ increased dramatically and was rapidly consumed after 3.2 min (Fig. 4C(vii)), which induced the generation of the product. Therefore, dynamic changes of the above intermediates involved in different potential pathways were provided for mechanism examination.

To further study the effect of OV on the selective C~N coupling, the free energy diagrams of the main route 1 → 3 → 4 → 5 → 7 on TiO₂ and TiO₂-OVs were calculated. The optimized structures of these intermediates were also determined (Fig. 4D). On TiO₂ (the black line in Fig. 4D), for generating 3 upon reaction between the adsorbed *o*-CAN (1) and intermediate 2, the energy change is -0.92 eV. Subsequently, 4 is obtained *via* intramolecular dehydrogenation of 3 with a potential barrier of 1.60 eV. Then, electrophilic substitution of -Cl *via* -



NH₂ can form intermediate 5 with the energy change of -1.75 eV. Finally, the product of 7-Cl-Phz (7) is obtained by further dehydrogenation of 5 through an exothermic process. While in the presence of OV, a lower Gibbs free energy for obtaining each intermediate is observed (the red line in Fig. 4D). In particular, the energy barrier of the step from intermediate 3 to intermediate 4 on TiO₂-OVs (1.03 eV) is lower than that of TiO₂ (1.60 eV), indicating that vacancies can reduce the energy barrier of this step. Therefore, except for promoting electron transfer, regulating molecular adsorption, modulating O₂ and substrate activation, the *in situ* generated OV can also lower the potential barriers of intermediate formation.

Mechanism of OV-controlled competitive photooxidation

On the basis of both experimental and theoretical examinations, the mechanism of the OV-controlled selective C–N coupling is proposed in three routes (Fig. 5). Initially, the substrate (1) could be adsorbed on the surface, resulting in weak coordinations *via* N–Ti or Cl–Ti. This allows electron injection from 1 to the conduction band (CB) of TiO₂ to give 1^{•+} (m/z 142). Notably (in Route I), the intermolecular hole transfer between 1^{•+} and neutral 1 would be initiated to afford 7-Cl-Phz (7) along with the *in situ* generation of OV. Simultaneously, the heteroatom-containing 7-Cl-Phz product can be easily adsorbed onto TiO₂ *via* surface coordination, acting as the sensitizer for the subsequent reactions. This dye-sensitization process would produce a new electron donor level above the VB of TiO₂ from the 2p orbital of Cl or N, facilitating broader visible-light absorption than the self-sensitization between *o*-CAN and TiO₂. Upon light irradiation, positive charge (h⁺) could be left at the heteroatoms after injecting electrons into the CB, which would ultimately transfer to O₂ to form O₂^{•-}. Subsequently, the intermolecular hole transfer between 7^{•+} and surface adsorbed 1 (mainly by N–Ti) would afford 1^{•+} and regenerate the surface-bound sensitizer 7, fulfilling the cyclic dye-sensitization of TiO₂.

Thereafter (in Route II), 1^{•+} would be oxidized by O₂^{•-} (obtained in Route I) to generate 2 (m/z 141)⁵⁴ and H₂O₂. Subsequently, 2 reacts with neutral 1 to generate 3 (m/z 283). Then, in the presence of H₂O₂, 3 undergoes intramolecular oxidative dehydrogenation to form 4 (m/z 281), followed by cyclization to form intermediate 5 (m/z 247). Further oxidation of 5 would afford the half-oxidized phenazine (Fig. 4A), which acts as the precursor of the product 7.⁵⁵ In addition (in Route III), 2 could also react with neutral 1 to generate intermediate 6 (m/z 249). Subsequently, 5 would be generated upon the oxidation and cyclization of 6, which supports the obtaining of product 7 as Route II indicated. Therefore, upon the absorption of the substrate and product on OV-containing TiO₂, intermolecular hole transfer would be initiated to facilitate the sensitization and step-by-step C–N coupling. This clearly reveals the mechanism of efficient and selective C–N coupling *via* photooxidation under visible light, even using small colorless organics as the reactant.

Conclusion

In conclusion, *in situ* generated OV on TiO₂ significantly facilitate the competitive visible-light photooxidation of colorless 4-chlorobenzene-1,2-diamine for selective C–N coupling rather than dechlorination degradation. By on-line MS examinations, the structural identification and dynamic monitoring of intermediates were successfully employed. This greatly supported the evaluation of interactions between the catalyst and reaction species or generated sensitizer, as well as examinations of energy changes upon molecular adsorption and transformation. The C–N coupling product was easily adsorbed on TiO₂, which extended the light absorption *via* surface coordination. The surface OV enhanced the interfacial charge transfer to promote selective C–N coupling *via* regulating the adsorption of the substrate and sensitizer, lowering the energy barrier and modulating O₂ activation on TiO₂. Based on this work, two general rules of visible light catalysis for photooxidation of small colorless organics could be drawn: (1) the synthetic product containing abundant heteroatoms can act as a better sensitizer than the substrate to exhibit higher reaction efficiency. (2) Oxidation of the substrates could be mediated by regulating the concentration of OV on the semiconductor catalyst. This work has initiated the comprehensive examination of both the catalyst and reaction species, which would promote the development of visible light catalysis.

Data availability

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

Author contributions

J.-H. S. and N. N. designed the project. J.-H. S. performed the experiments and wrote the manuscript. X.-Y. G. contributed to the theoretical calculations and the revision of the manuscript. Y.-X. G., M. Z. and Q. Z. participated in the characterizations of

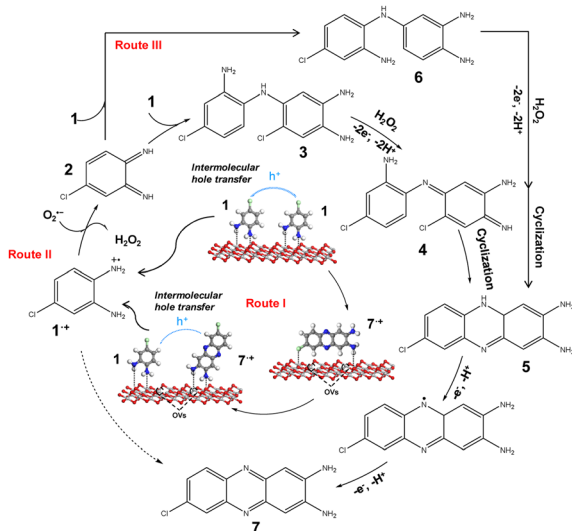


Fig. 5 Mechanism of OV-controlled selective C–N coupling.



the catalyst. G.-H. H., X.-N. W., Y.-Y. Y. and J. O. contributed to the mechanism studies. All authors have given approval to the final version of the manuscript. N. N. directed the research.

Conflicts of interest

There are no conflicts to declare.

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