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1 Introduction

Significant water contamination issues have been brought about by the widespread use and unreasonable discharge of organic compound products such as dyes, insecticides, and personal care items.¹⁻³ These organic pollutants pose a major hazard to human health, since they are not only extensively dispersed but also extremely poisonous. Photocatalytic technology is considered the most effective technique for environmental rehabilitation by eliminating the aforementioned organic contaminants owing to its simplicity of use, low cost, and absence of secondary contamination compared to conventional water treatment technologies (e.g. adsorption, flocculation, and advanced oxidation.4-8). However, the presence of diurnal alternation and the shading of sunlight by clouds under natural conditions severely limits the utilization of sunlight for semiconductor photocatalysis and makes the photocatalytic reaction time-limited in practical applications.9,10 Therefore, it is of great importance to find materials that can effectively store solar energy and continue catalytic

Correction and new understanding of the reactivity of illuminated tungsten trioxide in the dark: antecedents and consequences of photo-storage electrons triggering Fenton reactions[†]

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Water treatment via photocatalytic steps is severely constrained by the alternation of day and night and the shading of sunlight by clouds. As an energy storage photocatalyst, WO3 has been reported to function partially even under dark conditions, which is of great significance for the construction of round-theclock photocatalysts. However, pertinent reaction processes, especially the function of stored electrons in WO3 during catalytic reactions in the dark, have not been fully analyzed. Herein, we synthesized hexagonal WO₃ hydrate (h-WO₃·0.46H₂O) and used extensive experiments and theoretical calculations to investigate its structure, photochromic capability, and dark reaction activity. Due to the strong interaction between the protons from lattice water and bridging oxygen atoms (W-O-W), h-WO3.0.46H2O exhibited unexpected irreversible photochromic properties under visible light. Our study is the first to show that light-irradiated h-WO₃ \cdot 0.46H₂O can activate H₂O₂ to form \cdot O₂⁻, \cdot OH, and 1 O₂, which are responsible for the efficient degradation of organic pollutants, but it could not produce reactive oxygen species (ROS) via the 1e⁻/2e⁻ ORR in the dark. This study not only provides a novel and inexpensive Fenton catalyst for wastewater treatment, but also proposes a new approach for utilizing solar energy under dark conditions.

> reactions in the dark to achieve full utilization of solar energy and round-the-clock photocatalytic reactions.

> WO₃ is considered promising for the storage of solar energy and round-the-clock photocatalytic reactions owing to the reversible cycle of W⁶⁺/W⁵⁺. The reported mechanism can be represented by eqn (1) and (2), where M stands for H or alkali metals.11,12

 $WO_3 + xe^- (CB) + xM^+ \rightarrow M_x W^{5+}_{x} W^{6+}_{1-x}O_3 \text{ (charging)} (1)$

$$M_x W_x^{5+} W_{1-x}^{6+} O_3 \rightarrow WO_3 + xe^- + xM^+ \text{ (discharging)} \quad (2)$$

Based on the reversible structural transformation described above, many research studies on the degradation of organic pollutants by round-the-clock photocatalysts containing WO₃ have been reported (Table S1[†]). Although WO₃ exhibits obvious storage properties for photogenerated electrons and many studies have been reported on its properties and mechanisms for round-the-clock catalytic degradation of pollutants, there are still several issues that have not been fully addressed. Firstly, the limited storage capacity of photogenerated electrons and the rapid release of stored electrons lead to shorter catalytic reaction times under dark conditions. Secondly, low catalytic reaction activity under dark conditions cannot be matched with that under light illumination. Most importantly, the mechanism of catalytic reactions under dark conditions is ambiguous.

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For most reported dark reaction mechanisms of WO₃-based round-the-clock photocatalysts in the absence of light, the degradation of organic pollutants in the dark is attributed to the formation of \cdot OH (decomposition of H₂O₂ formed by the oxygen reduction reaction) or \cdot O₂⁻ through the oxygen reduction reaction (ORR) by stored electrons (Table S1†). It is worth noting that the electrons stored in WO₃ may reduce oxygen molecules under dark conditions through the following possible pathways (eqn (3)-(7)):¹³⁻¹⁵

$$O_2 + e_{CB}^- \rightarrow O_2^- \vec{E} = -0.33 V_{NHE}$$
 (3)

$$O_2^- + H^+ \rightarrow OOH$$
 (4)

OOH + H⁺ +
$$e_{CB}^- \rightarrow H_2O_2 E^\circ = +1.44 V_{NHE}$$
 (5)

$$O_2 + 2e_{CB}^- + 2H^+ \rightarrow H_2O_2 E^\circ = +0.68 V_{NHE}$$
 (6)

$$O_2 + 4e_{CB}^- + 4H^+ \rightarrow 2H_2O E^\circ = +1.23 V_{NHE}$$
 (7)

It is worth mentioning that WO_3 is a typical oxidation type semiconductor with a more positive conduction band (CB) position (>0 V_{NHE}).¹⁶⁻²⁰ So, the photogenerated electrons stored in the CB of WO₃ cannot reduce O₂ to O_2^- ($E^\circ = -0.33 V_{\text{NHE}}$) from a thermodynamic point of view, and therefore there is no way to produce H_2O_2 via eqn (4) and (5). Although the potential of the concerted two-electron oxygen reduction reaction (concerted 2e⁻ ORR, eqn (6)) seems to be appropriate, the fourelectron ORR to H_2O (eqn (7)) is thermodynamically more favourable because the electrode potential of the 4e⁻ ORR $(+1.23 V_{\text{NHE}})$ is more positive than that of the concerted $2e^{-}$ ORR $(+0.68 V_{\rm NHE})$. Obviously, in order to achieve the performance of WO₃ with photo-stored electrons to degrade pollutants in the dark matching that under light illumination, it is necessary to clarify the active species production and catalytic mechanism in the absence of light.

Herein, we synthesized a crystalline hexagonal WO₃ hydrate by a simple oil-bath heating method, which exhibited excellent and stable photogenerated electron storage capacity. In addition to the conversion of W(vI) to W(v) in hexagonal WO₃ hydrate under light illumination, the lattice water molecules stored in the intracrystalline tunnels undergo splitting. The generated protons will then interact strongly with the bridging oxygen atoms in the structure, changing the colour of the hexagonal WO₃ hydrate from white to blue, which can remain stable in the atmosphere for a long time (>24 h). In contrast to the results reported in the literature, we verified that the discoloured hexagonal WO3 hydrate could not reduce O2 molecules to reactive oxygen species (ROS) via the single-electron pathway or two-electron pathway. Interestingly, the electrons stored in the light-irradiated hexagonal WO₃ hydrate, however, can significantly activate H₂O₂ in the dark, generating ROS ($^{\circ}O_2^{-}$, $^{1}O_2$ and $^{\circ}OH$) for the efficient degradation of organic pollutants under dark conditions. When dark-Fenton reaction (DFR) is completed, the blue hydrated WO₃ can regain its original colour and then be recycled for the next catalytic reaction process.

2 Experimental

2.1. Materials

Sodium tungstate dihydrate (Na₂WO₄·2H₂O) was obtained from Energy Chemical Reagent Co., Ltd. Concentrated hydrochloric acid (37%, HCl) and polyvinylpyrrolidone with a molecular weight of 24 000 (PVP, M = 24 K) were purchased from Tianjin Kermel Chemical Reagent Co., Ltd .2 *sec*-Butyl-4,6dinitrophenol (DNBP) was obtained from Aladdin Reagent Co., Ltd In addition, deionized water was used in all the operations. All reagents were used as received without further purification.

2.2. Synthesis of samples

2.2.1. Synthesis of hexagonal WO₃ hydrate. The hexagonal WO₃ hydrate was synthesized by a simple oil-bath heating method. Typically, 2 g of Na₂WO₄ · 2H₂O was added to a 100 ml beaker with 30 ml of deionized water and dissolved under continuous magnetic stirring. Then, 10 ml of aqueous solution containing 15 mg of PVP was added dropwise to the above solution and kept stirring for 5 min. After that, 1.5 mmol Na₂SO₄ was poured into the above transparent solution and stirred for 30 min to dissolve it. The system was then heated to 90 °C in an oil bath and kept warm for 10 min. Subsequently, about 4.5 ml of 3.0 M HCl solution was added dropwise to the reaction was continued at 90 °C for 3 h. Finally, the obtained white product was washed 4 times with deionized water, freeze-dried, and named h-WO₃ · xH_2O .

2.2.2. Synthesis of illuminated hexagonal WO₃ hydrate. 80 mg of h-WO₃ hydrate was dispersed into 30 ml of deionized water and kept stirring for *t* min under visible light irradiation ($\lambda > 420$ nm). Upon illumination, the colour of h-WO₃ hydrate gradually changed from white to blue. Then, the obtained blue precipitate was collected after centrifugation, washing and lyophilization. The final light irradiated product was named LP*t* min, where *t* represents different light irradiation times.

2.3. Characterization

A JEM-F200 field emission transmission electron microscope (TEM) and Rigaku D/Max 2400 diffractometer equipped with monochromatized Cu K α radiation (l = 0.15406 nm) were used to investigate the morphology and structures of the materials. Fourier transform infrared (FTIR) spectra were obtained with a Thermo Nicolet Avatar 360 Fourier transform infrared spectrometer. The chemical state of the elements was determined using a Thermo Fisher ESCALAB Xi⁺ X-ray photoelectron spectrometer (XPS) equipped with a scanning range of 100-4000 eV, a 180° double-focusing hemispherical analyzer with a 128channel detector, and an Al Ka micro-focused X-ray source. The optical absorption performances and optical band gap of the samples were obtained using a HITACHI U-3900 spectrometer, and BaSO₄ served as a standard reference substance. Electron paramagnetic resonance spectroscopy (EPR) was recorded on a Bruker A200 spectrometer at room temperature. The internal structure of the samples was investigated using a confocal Raman spectrometer (Raman, LabRAM HR Evolution, Horiba, France) with the scanning range of 100–1000 cm⁻¹. HP6890GC/ 5973MSD apparatus was employed to analyze intermediate products in the degradation of the target pollutant DNBP by gas chromatography-mass spectrometry (GC-MS) mode.

Additionally, temperature programmed desorption (TPD) curves were recorded on a TGA8000-Frontier-Clarus SQ8T instrument at a heating rate of 10 °C min⁻¹ in Ar. The heating range is room temperature-700 °C. The mass loss of the sample above 100 °C was ascribed to the wastage of the lattice water molecules.²¹ So, the amount of lattice H₂O per WO₃ formula unit (*x*) can be calculated by eqn (8):

$$x = n_{\rm H_2O}/n_{\rm WO_3} = (m_0 A_2/A_1 M_{\rm WO_3})/(m_0 (A_1 - A_2)/A_1 M_{\rm H_2O})$$

= [(A_1 - A_2)M_{\rm WO_3}] (8)

wherein $n_{\rm H_2O}$, $n_{\rm WO_3}$, m_0 , A_1 , A_2 , $M_{\rm WO_3}$ and $M_{\rm H_2O}$ refer to the number moles of lattice H₂O molecules present in a certain mass of WO₃·*x*H₂O (mol), the number moles of WO₃ present in a certain mass of WO₃·*x*H₂O (mol), a certain mass of WO₃·*x*H₂O (g), the mass ratio of the sample above 100 °C (%), the mass ratio of the sample above the temperature (400–550 °C) at a stable mass (%), the molar mass of WO₃ (g mol⁻¹) and the molar mass of H₂O (g mol⁻¹), respectively.

2.4. Quantitative experiments on stored electrons in LPt min samples and the oxidation experiments of I^- for different systems

The detailed procedures are shown in the ESI (Text S1.1 and Text S1.2). \dagger

2.5. Theoretical calculations

Information on theoretical calculations is shown in ESI (Text S1.3).†

2.6. Performance evaluation of catalysts

The removal experiments of pollutants by the synthesized samples were carried out on a CEL-LAB500 photoreactor (CEAULIGHT Co., Ltd) equipped with a 500 W Xe lamp with a 420 nm cut-off filter. The average light intensity over the duration of each experiment was measured to be 80 mW cm⁻². DNBP, a typical dinitrophenol compound which is widely used in the petrochemical industry and agriculture, was chosen as a target organic contaminant. Typically, 30 mg of the assynthesized catalyst was dispersed into 30 ml of the target pollutant solution (20 mg L^{-1}) in a quartz tube. Prior to turning on the light, the suspension was stirred in the darkness for 30 min to establish adsorption-desorption equilibrium. The system was then subjected to visible light irradiation under magnetic stirring. Samples of 3 ml were withdrawn at different time intervals, and centrifuged and the supernatant was analysed for evolution in concentration of DNBP spectrophotometrically at λ_{max} of 376 nm. The dark Fenton-like reaction (DFR) of the samples was triggered after photocatalytic experiments in the same setup. After turning off the light, 150 µL of H_2O_2 solution (10 wt%) was added into the quartz tube and kept stirring. Afterwards, the variation of DNBP concentration was

monitored by the same method and procedure as described above for photocatalysis measurements. In particular, the DFR process was performed under completely dark conditions.

2.7. Electrochemical measurements

All electrochemical measurements were performed on an electrochemical workstation (Chenhua CHI 660E, Shanghai Chenhua Instrument Co., Ltd) equipped with a standard threeelectrode system. Mott–Schottky (MS) measurements were performed with the same setup at a frequency of 1000/2000/3000 Hz with an amplitude of 5 mV. Linear sweep voltammetry (LSV) was performed through a rotating ring disk electrode (RRDE) with a Pt ring electrode in O₂-saturated solution, and the working electrode was prepared by loading the catalyst on a glassy carbon electrode. The number of electrons (*n*) transferred during the dark reaction can be calculated using the following equation:

$$n = (4 \times I_{\rm D})/(I_{\rm D} + I_{\rm R}/N)$$
 (9)

wherein $I_{\rm D}$, $I_{\rm R}$ and *N* refer to the disk current (mA), the ring current (mA) and the collection efficiency (0.37), respectively. In addition, the number (*n*) of electron transfer involved in the overall O₂ reduction was also calculated from the linear regression of the Koutecky–Levich (K–L) plot using the following equations (eqn (10) and (11)):

$$1/j = 1/j_{\rm d} + 1/j_{\rm k} = B^{-1}\omega^{-1/2} + j_{\rm k}^{-1}$$
(10)

$$B = 0.2nFv^{-1/6}CD^{2/3} \tag{11}$$

where *j* is the current density (mA cm⁻²), j_d is the diffusionlimited current, j_k is the kinetic current density, ω is the angular velocity of the disk electrode (rpm), ν is the kinetic viscosity of water (0.01 cm² s⁻¹), *C* is the concentration of dissolved O₂ in water (1.26 × 10⁻³ mol L⁻¹), and *D* is the diffusion coefficient of dissolved O₂ (2.7 × 10⁻⁵ cm² s⁻¹).

All measurements were performed with the same setup in 30 ml Na₂SO₄ solution (0.5 M). The experiments were carried out at ambient temperature (25.0 \pm 0.5 °C).

3 Results and discussion

The apparent morphology and microstructure of the samples were further characterized by SEM and TEM measurements (Fig. S1[†] and 1a). It can be clearly seen that the h-WO₃ hydrate sample shows a nano-strip morphology with a length of ~100 nm and a width of ~20 nm. The fast Fourier transform (FFT) of the HRTEM image with the lattice plane intensity profile (inset of Fig. 1b) exhibited an interplanar spacing of 0.39 nm clearly, which can be ascribed to the (001) plane of hexagonal WO₃. Additionally, the results from the selected area electron diffraction (SAED) technique (Fig. 1c) showed that the h-WO₃ hydrate sample had a splendid crystal structure with planes (100), (002), (200), and (202) of hexagonal WO₃. As shown in Fig. 1d–g, the surface of the h-WO₃ hydrate sample is uniformly distributed with O, Na and W, wherein the Na

element comes from Na₂WO₄·2H₂O. On the side, the crystallographic information of the sample was obtained through powder X-ray diffraction (PXRD) analysis. The main peaks were observed at 13.96°, 22.72°, 24.33°, 26.85°, 28.17°, 33.58°, 36.57°, 49.96°, 55.33° and 55.51° (Fig. 1h), which, respectively, corresponded to Bragg reflections from the (100), (001), (110), (101), (200), (111), (201), (220), (202) and (221) crystalline planes of hexagonal WO₃ (space group P6/mmm),^{22,23} which was consistent with the SAED findings. Thermogravimetric analysis (TGA) revealed that the mass loss of the sample above 100 °C was about 3.47% (Fig. 1i), indicating the presence of $0.46H_2O$ (x = 0.46) per WO₃ formula unit (calculation details are shown in Text 2.3). Based on the above results, the crystallographic structure of the prepared h-WO₃ · 0.46H₂O was refined (Fig. 1h), and the Rietveld refinement data are summarized in Table S2.† In h-WO₃·0.46H₂O, the lattice water molecules are in the dodecagon tunnels along the z axis and reside between W-O layers along the z axis (Fig. 1j, S2a and b[†]).

The photochromic response of $h-WO_3 \cdot 0.46H_2O$ was investigated under continuous visible light illumination using a 500 W

Xe lamp equipped with a filter (>420 nm) for different irradiation times. As shown in Fig. S3,[†] h-WO₃·0.46H₂O gradually changed from milky white to dark blue with the increase of illumination time. The photochromic process proceeded slowly and the color of the sample hardly changed significantly after 80 min of irradiation. This phenomenon can also be confirmed by the absorbance and reflectance spectra of different illuminated h-WO₃·0.46H₂O samples (Fig. 2a and S4[†]). The absorbance and reflectance of visible light area for h-WO3 · 0.46H2O underwent a significant enhancement with the increase of illumination time. Compared with the sample LP-80 min, the absorbance and reflectance of the sample LP-100 min were hardly enhanced significantly, indicating that the photochromism of h-WO₃·0.46H₂O had reached saturation after visible light irradiation for about 80 min. Moreover, the illuminated h-WO₃·0.46H₂O exhibited a surprising stability in air, with the absorbance of the sample LP-80 min at 700 nm decreasing by only 12% after more than 24 h (Fig. S5[†]), reflecting the excellent stability of the stored photogenerated electrons. The stored electrons in illuminated h-WO₃·0.46H₂O were quantified by the



Fig. 1 (a) TEM image and (b) HR-TEM image (the inset shows the corresponding fast Fourier transform (FFT) patterns and the lattice plane intensity profile). (c) SAED patterns of $h-WO_3 \cdot 0.46H_2O$. (d) The annular dark field (ADF) image and (e–g) elemental mapping images of $h-WO_3 \cdot 0.46H_2O$. (h) XRD pattern and the Rietveld refinement of $h-WO_3 \cdot 0.46H_2O$. (i) TG curve of unilluminated $h-WO_3 \cdot 0.46H_2O$. (j) The refined crystal structure unit of $h-WO_3 \cdot 0.46H_2O$ viewed toward the *xy* plane.

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Fig. 2 (a) Absorbance spectra of different illuminated $h-WO_3 \cdot 0.46H_2O$ samples. (b) The reduction experiments of KMnO₄ using different catalysts. The reduction experiments of KMnO₄ with initial concentrations of (c) 120 ppm, (d) 150 ppm, and (e) 200 ppm using the LP-80 min catalyst. (f) The concentration of stored electrons for different illuminated $h-WO_3 \cdot 0.46H_2O$ samples. (g) TG curve of unilluminated $h-WO_3 \cdot 0.46H_2O$. (h) The XRD spectra and (i and k) partial enlarged view of XRD patterns for different illuminated $h-WO_3 \cdot 0.46H_2O$. (j and l) Schematic diagram of lattice changes for $h-WO_3 \cdot 0.46H_2O$ under illumination.

reduction experiments of KMnO4 (the experimental details are displayed in Text S1.1[†]), in view of the good linearity between KMnO₄ solution concentration and absorbance (Fig. S6[†]). Based on the Lambert-Beer Law and the stoichiometric relationships of chemical reaction $(MnO_4^{-} + 5W(v) + 8H^{+} - Mn^{2+})$ $+5W(v_1) + 4H_2O$, Fig. S7[†]), the concentration of stored electrons for LP-80 min can be calculated to be 0.6033 mmol g^{-1} by comparing the absorbance changes of KMnO₄ solution before and after the reduction reaction (Fig. 2b). The reliability of the result was also validated by the reduction experiments of KMnO₄ solution with different initial concentrations (Fig. 2c-e). Similarly, the concentration of stored electrons for different LPt min can also be obtained (Fig. 2f). Obviously, when illuminated for more than 80 min, the storage capacity of h-WO₃·0.46H₂O for photogenerated electrons is close to saturation. The above results revealed the slow rate of coloration of h-WO3 · 0.46H2O and the anomalous stability of the illuminated h- $WO_3 \cdot 0.46H_2O$, which also imply that the photochromic mechanism of h-WO₃ has its own peculiarities.

TGA of illuminated h-WO₃ \cdot 0.46H₂O and XRD measurements of h-WO₃ \cdot 0.46H₂O with different illumination times were performed to investigate the mass change and phase evolution of the sample during the photochromic process. As shown in Fig. 2g, the mass attenuation in the range of >100 °C was 3.41%, which is very close to that of the unilluminated hWO₃·0.46H₂O, reflecting that no significant loss of lattice water of h-WO₃·0.46H₂O occurred before and after illumination. In addition, the XRD diffraction peaks of h-WO3 · 0.46H2O were well preserved after illumination (Fig. 2h), indicating that irradiation only changed its color but did not affect its crystal phase. However, changes in the position of the diffraction peaks can still be observed from the partial magnified view of the XRD patterns. During illumination, the (0 0 1) diffraction peak of h-WO₃·0.46H₂O gradually shifted to the high angle direction with the increase of time (Fig. 2i), reflecting the structural contraction of the crystal lattice along the z axis that occurred in h- $WO_3 \cdot 0.46H_2O$ (Fig. 2j), which may be caused by the strong interaction between the protons originating from the lattice water and the bridging oxygen atoms during the formation of H_xWO_3 . Also, the (1 0 0) peak shifts to the left (Fig. 2k), suggesting the elongation of the crystal along the xy plane (Fig. 2l), which also indicates the existence of such strong interaction. The lattice elongation along the xy plane and crystal contraction along the z axis facilitate the entry of water molecules into the lattice of h-WO₃·0.46H₂O from outside to replenish the lattice water molecules consumed by the photochromic effect.

To further clarify the particular photochromic mechanism of $h-WO_3 \cdot 0.46H_2O$, additional spectroscopic tests were used to analyse the surface electronic state and chemical structure of $h-WO_3 \cdot 0.46H_2O$ before and after light irradiation. Full XPS



Fig. 3 (a) XPS survey spectra of $h-WO_3 \cdot 0.46H_2O$ before and after illumination. High resolution XPS spectra of (b) O 1s and (c) W 4f in unilluminated $h-WO_3 \cdot 0.46H_2O$ and illuminated $h-WO_3 \cdot 0.46H_2O$. (d) The EPR profiles and (e) Raman spectra of $h-WO_3 \cdot 0.46H_2O$ before and after illumination. (f) The partial enlarged view of Raman patterns of the samples. (g) The FT-IR spectra and (h) partial enlarged view of FT-IR spectra for $h-WO_3 \cdot 0.46H_2O$ with different illumination times. (i) Schematic illustration of photochromism process for $h-WO_3 \cdot 0.46H_2O$.

spectra (Fig. 3a) showed that W and O are the major elements in the samples without other impurity elements. In addition, the high resolution XPS model was performed to study the chemical state of O and W elements. After convolution of the XPS spectra, both O 1s and W 4f spectra exhibit two individual peaks. The peaks at 530.49 eV and 532.34 eV in the O 1s spectrum of the unilluminated h-WO₃·0.46H₂O are assigned to the lattice O in hexagonal WO₃ and the adsorbed water molecules on the surface of the sample (Fig. 3b),²⁴⁻²⁷ respectively. As for the W 4f spectrum of the unilluminated h-WO₃·0.46H₂O, the two peaks at 35.79 eV and 37.93 eV are attributed to the spin-orbit splitting of the W 4f components (W $4f_{7/2}$ and W $4f_{5/2}$) (Fig. 3c).²⁸⁻³⁰ After visible light irradiation, both the O 1s and W 4f peaks of h-WO₃·0.46H₂O shifted toward the lower binding energy, indicating that the photochromic process of h-WO₃·0.46H₂O was accompanied by an increase in the local electron density of O and W.³¹ The electron paramagnetic resonance (EPR) spectra (Fig. 3d) of the illuminated h-WO₃·0.46H₂O showed a sharp signal at about 3600 G with g = 1.792 compared to the unilluminated h-WO₃ \cdot 0.46H₂O, reflecting the existence of W⁵⁺, ^{32,33} which can be attributed to the increase in the electron cloud density of W due to the excitation of visible light and coincides with the XPS results.

Raman and FT-IR spectra provide detailed information on the surface functional groups and chemical bonds of the samples to analyze the changes of $h-WO_3 \cdot 0.46H_2O$ before and after light illumination. From the results of Raman measurements, the peaks assigned to the W-O-W bending modes/the bending vibration δ (W⁶⁺-O) (247 cm⁻¹, 323 cm⁻¹),³⁴ O-W-O stretching modes (679 cm⁻¹, 807 cm⁻¹)³⁴⁻³⁶ and the stretching vibration of terminal W=O bonds (946 cm⁻¹)^{34,35} of h-WO3 · 0.46H2O were substantially attenuated after illumination (Fig. 3e), suggesting that the chemical bond between W and O in h-WO₃·0.46H₂O was seriously weakened by the strong interaction between the protons originating from lattice water and the bridging oxygen atoms. In addition, the partial enlarged view of Raman patterns of the samples (Fig. 3f) showed that the dominant peaks of illuminated h-WO3 · 0.46H2O shifted toward the lower wavenumber compared to the unilluminated sample, also verifying the change in Raman vibrational energy level due to the interaction between the protons from lattice water and the bridging oxygen atoms. As shown in Fig. 3g, all the samples exhibited similar IR absorption spectra including the main IR absorption peaks at 3430 cm^{-1} (the stretching vibrations of surface absorbed water),^{31,35} 2380 cm⁻¹ (asymmetric stretching vibrations of surface absorbed CO_2),³⁷ 1610 cm⁻¹ (the bending vibrations of surface absorbed water)31,38 and 1100-500 cm⁻¹ (the typical WO₆ vibration bands include the O-W-O band and W=O band).^{34,35} Noteworthily, the bending vibration peak of surface absorbed water for illuminated h-WO3 · 0.46H2O showed an obvious red shift compared to unilluminated h- $WO_3 \cdot 0.46H_2O$, which can be ascribed to the compensation

effect of external water molecules on lattice water molecules. Additionally, the vibrational absorption of O–W–O and W==O for h-WO₃ \cdot 0.46H₂O decreased gradually with increasing irradiation time (Fig. 3h). The results are consistent with the results of XRD and Raman measurements.

From the results of the above series of spectroscopic tests, the stable photochromic process of h-WO₃·0.46H₂O can be reasonably explained (the schematic illustration is shown in Fig. 3i): under visible light irradiation, the local electronic density of h-WO₃·0.46H₂O increases and lattice water located in the dodecagon tunnels along the *z*-axis of h-WO₃ is decomposed. The generated protons then interact with the bridging oxygen atoms in the WO₆ structure, leading to the weakening of O–W–O as well as W=O bonds. Meanwhile, the interaction also induces a distortion of the h-WO₃·0.46H₂O lattice (lattice relaxation along the *xy* plane and contraction along the *z* axis), which helps the external water molecules to replenish the lattice water molecules consumed by the photochromic effect. It is due to this special photochromic mechanism that the illuminated h-WO₃·0.46H₂O can maintain its color stability in air for a long time.

In our study, the blue color of the illuminated h-WO₃·0.46H₂O does not easily fade in air, it can be decolorized

by the oxidation of strong oxidizing agents (e.g., H_2O_2), and the oxidation experiments of I⁻ (the experimental details are shown in Text S1.2[†]) were carried out to monitor the consumption of H_2O_2 . As shown in Fig. 4a, the absorbance of I_3^- is very close only in the H₂O₂ and unilluminated h-WO₃·0.46H₂O/H₂O₂ systems, indicating that the unirradiated h-WO₃·0.46H₂O did not react significantly with H₂O₂. As for the illuminated h-WO₃·0.46H₂O sample, its blue colour gradually fades after the addition of H₂O₂ (inset in Fig. 4a), reflecting the gradual conversion of the W(v) species formed through light illumination to W(vi) species. Interestingly, the characteristic absorbance of I3- in illuminated h-WO3·0.46H2O/H2O2 did not decrease but exhibited a significant increase compared to that in the H₂O₂ or unilluminated h-WO₃·0.46H₂O/H₂O₂ systems. Besides, Fig. 4b showed that illuminated h-WO₃·0.46H₂O has a considerably greater H₂O₂ reduction current in the dark compared to unilluminated h-WO₃·0.46H₂O, and the H₂O₂ reduction current on illuminated h-WO3.0.46H2O increased with increasing the addition amount of 10 wt% H₂O₂ solution from 30 to 150 μ L (Fig. 4c). This proves that there is a unique electron transfer between illuminated h-WO3 · 0.46H2O and H_2O_2 from the former to the latter. The above surprising results



Fig. 4 (a) The oxidation experiments of I⁻ for different catalysts. (b) LSV curves of the electrode of unilluminated h-WO₃ \cdot 0.46H₂O and illuminated h-WO₃ \cdot 0.46H₂O in the presence of 150 μ L H₂O₂ (10 wt%) in 30 ml N₂-saturated 0.5 M NaNO₃ solution under dark conditions. (c) LSV curves of the illuminated h-WO₃ \cdot 0.46H₂O electrode in 30 ml N₂-saturated 0.5 M NaNO₃ solution containing different [H₂O₂] under dark conditions. MD simulation for (d) unilluminated h-WO₃ \cdot 0.46H₂O/H₂O₂ and (e) illuminated h-WO₃ \cdot 0.46H₂O/H₂O₂ systems.

suggest that the illuminated h-WO₃ \cdot 0.46H₂O does not undergo the expected oxidation process in the presence of H₂O₂, but rather exhibits the activity of Fenton reaction, resulting in more efficient oxidation of I⁻ by generating more reactive oxygen species (ROS).

To further comprehend the process underlying the activation of H_2O_2 molecules by the illuminated h-WO₃·0.46H₂O, the H_2O_2 adsorption on the (001) surface of the h-WO₃·0.46H₂O before and after illumination was compared by theoretical calculations (Text S1.3†). From the molecular dynamics (MD) simulations, the H_2O_2 molecule with one side protruding toward the vacuum binds to the (001) surface of unilluminated h-WO₃·0.46H₂O (Fig. 4d), which is clearly not conducive to adequate contact between the O–O bond and the catalyst surface. Contrarily, the H_2O_2 molecule can be permanently adsorbed horizontally on the (001) surface of illuminated h- $WO_3 \cdot 0.46H_2O$ (Fig. 4e). The favorable contact between the H_2O_2 molecule and illuminated h- $WO_3 \cdot 0.46H_2O$ (001) surface creates the conditions for the activation step of the O–O bond.

In addition, the H_2O_2 adsorption on the (001) surface of the h-WO₃·0.46H₂O before and after illumination was also analysed from a thermodynamic perspective. As presented in Fig. 5a, the E_a of H_2O_2 for illuminated h-WO₃·0.46H₂O is -2.89 eV, much more negative than that of the unilluminated sample (-1.57 eV), which means that the H_2O_2 molecule is more easily adsorbed on the surface of illuminated h-WO₃·0.46H₂O than unilluminated h-WO₃·0.46H₂O.³⁹ Moreover, the longer length (3.145 Å, Fig. 5b) of O–O bond of H_2O_2 molecule adsorbed on illuminated h-



Fig. 5 (a) E_a (adsorption energy) of H_2O_2 on the (0 0 1) surfaces of unilluminated and illuminated h- $WO_3 \cdot 0.46H_2O$. The variations in O–O bond length of the H_2O_2 molecule before and after its adsorption on the (0 0 1) surface of (b) unilluminated and (c) illuminated h- $WO_3 \cdot 0.46H_2O$. (d) The comparison of concentration of DNBP degradation in h- $WO_3 \cdot 0.46H_2O$ under different conditions. Degradation of DNBP with illuminated h- $WO_3 \cdot 0.46H_2O$ is under different conditions. Degradation of DNBP with illuminated h- $WO_3 \cdot 0.46H_2O$ via the DFR process under different (e) amounts of H_2O_2 solution (10 wt%) and (f) pH values. (g) Mott–Schottky plots at different testing frequencies for unilluminated h- $WO_3 \cdot 0.46H_2O$. (h) RRDE polarization curves over the illuminated h- $WO_3 \cdot 0.46H_2O$ electrode at 1600 rpm in O_2 -saturated electrolyte using the ring current and the disk current. (i) Koutecky–Levich (K–L) plots obtained *via* RDE analysis in an O_2 -saturated solution at -1.0 V (vs. Ag/AgCl) under the dark condition.



Fig. 6 (a) Trapping experiments of reactive species with illuminated $h-WO_3 \cdot 0.46H_2O/H_2O_2$ in the dark. (b–d) EPR signals of 'OH, 'O₂⁻, and ¹O₂ in the presence of DMPO, DMPO, and TEMP among different reaction systems, respectively. (e) The adsorption energy diagrams for HOO', ¹O₂ and 'OH on the (0 0 1) surface of illuminated $h-WO_3 \cdot 0.46H_2O$. (f) The reusability of $h-WO_3 \cdot 0.46H_2O$ for degradation of DNBP *via* dark adsorption-photodegradation-DFR process. (g) The concentration of stored electrons for LP-80 min through different cycle times. (h) Schematic illustration of the photochromic mechanism and DFR mechanism for $h-WO_3 \cdot 0.46H_2O$.

 $WO_3 \cdot 0.46H_2O$ compared with unilluminated h-WO₃ $\cdot 0.46H_2O$ (1.937 Å, Fig. 5c) indicates the activation tendency. Adequate contact between the H_2O_2 molecule and illuminated h- $WO_3 \cdot 0.46H_2O$ (001) surface can facilitate the activation of the O–O bond. Based on the above results of theoretical calculations, it can be easily drawn that the enhanced thermodynamic and kinetic ability of h-WO₃ $\cdot 0.46H_2O$ to adsorb H_2O_2 molecules after illumination creates the conditions for the initiation of the ensuing Fenton reaction.

DNBP was employed as the target pollutant to better illustrate the dark-Fenton reaction (DFR) activity of illuminated h-WO₃·0.46H₂O (Fig. 5d and Text S2.6[†]). Under illumination, the DNBP degradation performance of h-WO₃ · 0.46H₂O was so weak and just \sim 7% of DNBP was removed in 2 h. After turning off the light, the concentration of DNBP in illuminated h-WO₃·0.46H₂O remained almost constant, which indicated that the stored electrons in illuminated h-WO3 · 0.46H2O cannot induce the degradation of the target pollutant. However, more than 80% of DNBP in illuminated h-WO₃ · 0.46H₂O was removed in 1 h after adding H₂O₂, which far exceeded the activity of only H_2O_2 (11%) and unilluminated h-WO₃·0.46H₂O/H₂O₂ (24%) under the same conditions, reflecting the activation of H₂O₂ by illuminated h-WO₃·0.46H₂O. Furthermore, optimization of reaction conditions (Fig. 5e and f) and TOC removal amount (Fig. S8[†]) were explored. To further clarify the role of the stored

electrons in illuminated h-WO₃·0.46H₂O, a series of electrochemical tests were carried out. As shown in Fig. 5g, the Mott-Schottky curve at different testing frequencies exhibited a positive slope (n-type characteristic^{40,41}) and a flat band potential of 0.17 V vs. NHE; therefore, the conduction band edge (E_{CB}) of h-WO₃ \cdot 0.46H₂O is 0.17 V vs. NHE.^{42,43} Owing to the positive E_{CB} , illuminated h-WO₃·0.46H₂O cannot reduce O₂ by the singleelectron pathway (eqn (3)). Besides that, the reduction disk current (I_D) and the oxidation ring current (I_R) under different potential of the illuminated h-WO₃·0.46H₂O electrode in O₂saturated solution are shown in Fig. 5h. The invisible $I_{\rm R}$ and distinct $I_{\rm D}$ can be observed since the storage electrons in illuminated h-WO₃·0.46H₂O were constantly consumed by the O₂ molecule, and the average electron transfer number on illuminated h-WO₃ \cdot 0.46H₂O can be estimated to be ~4 by the formula (Text 2.7). The linear regression result of the Koutecky-Levich (K-L) plot showed that the number of electron transfer involved in the overall O₂ reduction was 3.98 (Fig. 5i and Text 2.7), which was in good agreement with the RRDE result. So, the illuminated h-WO3 · 0.46H2O in aqueous solution in the dark induces the four-electron ORR pathway (eqn (7)) to produce H₂O rather than the single-electron (eqn (3)) or two-electron pathway (eqn (6)) to generate H_2O_2 , which thermodynamically demonstrates that the stored electrons in illuminated h-WO₃ · 0.46H₂O cannot generate ROS via the ORR pathway in the dark.

The type of active species is important for understanding the mechanism of the DFR process; trapping experiments of reactive species among different systems in the dark are carried out to explore the contribution of different active species. Fig. 6a displayed that the degradation of DNBP was obviously suppressed after introducing high purity nitrogen or NaN₃ or isopropanol, indicating that 'O2⁻, 1O2 and 'OH were the active species in the process of DNBP degradation with the illuminated $h\text{-WO}_3 \cdot 0.46 H_2 O/H_2 O_2$ system in the dark. It can be also observed from the figure that the O_2^- is the prominent active species in the DFR process and the role of ${}^{1}O_{2}$ is also important. Furthermore, EPR measurements can also clearly show the kind of reactive species produced during the process of Fenton reaction. As shown in Fig. 6b, the illuminated h-WO₃·0.46H₂O/ H₂O₂ system exhibits the apparently stronger signal of 'OH than the unilluminated h-WO3 · 0.46H2O/H2O2 system and illuminated h-WO₃·0.46H₂O, which not only indicates that the electrons stored in illuminated h-WO₃·0.46H₂O cannot contribute to the degradation of the target pollutant directly, but also demonstrates its excellent activation capacity of H2O2. Similar results can be seen from the EPR spectra of $DMPO-O_2^-$ and TEMP-¹O₂ (Fig. 6c and d).

To better understand the process by which reactive species are produced on the catalyst surface, the adsorption energy of HOO' (since ' O_2^- is a Brønsted base with $pK_a = 9.12$,^{44,45} the

protonated form is used in the process of theoretical calculation), ${}^{1}O_{2}$ and ${}^{\cdot}OH$ on the illuminated h-WO₃ \cdot 0.46H₂O (0 0 1) surface was calculated by the DFT method, respectively. As shown in Fig. 6e, the values of E_{ads} for HOO', ${}^{1}O_{2}$ and 'OH are -0.70 eV, -2.49 eV and -3.48 eV, respectively. The smaller value of E_{ads} indicates that the adsorption between the adsorbent and adsorbate is unstable and adsorbate molecules are easily desorbed from the surface of the adsorbent.⁴⁶ Apparently, HOO' is more likely than ¹O₂ and 'OH to leave the surface of illuminated h-WO₃·0.46H₂O and participate in the degradation reaction of DNBP. In addition, if the dominant active species directly produced by the illuminated h-WO₃·0.46H₂O/H₂O₂ system is ${}^{1}O_{2}$ or 'OH, these two ROS will oxidize W(v) to W(v) because they are difficult to desorb from the surface of illuminated h-WO₃·0.46H₂O, which will cause the activity of the catalyst to decay and is inconsistent with experimental results. It can be concluded that the directly active species for the activation of H_2O_2 with illuminated h-WO₃·0.46H₂O is 'O₂⁻, and the formation of ¹O₂ and [•]OH is most likely the result of a chain reaction caused by O_2^- . In light of this, the generation of O_2^- , OH and 1O_2 in the illuminated h-WO₃ $\cdot 0.46H_2O/H_2O_2$ system in the dark can be demonstrated by the routes below (eqn (12)-(16)):47-50

$$[W_VO_6] + H_2O_2 \rightleftharpoons [O_6W_V(O_2H_2)]$$
(12)

9 9 9	Atom	Fuku	i (0) Fu		ui (-)	Fukui (+) e	
		Mulliken	Hirshfeld	Mulliken	Hirshfeld	Mulliken	Hirshfeld
	C (1)	0.048	0.061	0.023	0.043	0.074	0.078
0	C (2)	0.047	0.051	0.020	0.023	0.074	0.079
	C (3)	0.012	0.041	0.023	0.046	0.001	0.035
	C (4)	0.070	0.076	0.047	0.058	0.093	0.094
	C (5)	0.050	0.053	0.019	0.033	0.061	0.072
	C (6)	0.043	0.049	0.065	0.067	0.022	0.031
	C (7)	0.073	0.078	0.069	0.072	-0.028	0.006
	C (8)	-0.014	0.009	-0.013	0.007	-0.016	0.011
	C (9)	-0.014	0.007	-0.013	0.005	-0.016	0.008
-4.19xe ²	C (10)	-0.009	0.004	-0.007	0.004	-0.011	0.005
	O (11)	0.047	0.052	0.069	0.075	0.025	0.030
	N (12)	0.056	0.061	0.079	0.083	0.033	0.039
	N (13)	0.052	0.065	0.063	0.071	0.021	0.019
	O (14)	0.049	0.049	0.052	0.051	0.046	0.046
	O (15)	0.041	0.058	0.047	0.053	0.035	0.063
	O (16)	0.053	0.060	0.058	0.065	0.028	0.035
4.19×e ⁻²	0 (17)	0.035	0.041	0.059	0.066	0.030	0.036

Fig. 7 (a) The optimized molecular structure, (b) HOMO (-0.269 eV), (c) LUMO (-0.118 eV), and (d) electrostatic surface potential (ESP) of DNBP. (e) Fukui indices (e Å⁻³) of the DNBP molecule.

$$[O_6W_V(O_2H_2)] + H_2O_2 \rightarrow [W_{VI}O_6] + O_2^- + 2H_2O$$
(13)

$$O_2^- + H_2O_2 \rightarrow OH + O_2 + OH^-$$
, Haber–Weiss reaction,

$$\Delta G = -0.64 \text{ eV} = -62 \text{ kJ mol}^{-1}$$
(14)

$$2 \cdot O_2^- + 2H_2O \rightarrow {}^1O_2 + H_2O_2 + 2OH^-,$$

 $\Delta G = -0.65 \text{ eV} = -63.3 \text{ kJ mol}^{-1}$ (15)

$$O_2^- + OH \rightarrow O_2^+ + OH^-, \Delta G = -1.31 \text{ eV} = -127 \text{ kJ mol}^{-1}(16)$$

The reusability and stability of the catalyst were evaluated by 10 cycles of experiment. After 10 cycles, the DNBP degradation efficiency decreased from 92.3% to 75.6% (Fig. 6f), accompanied by the TOC removal also decreasing from 82.1% to 62.6% (Fig. S9a†), which can be ascribed to the reduction in the number of ROS generated from the DFR process with increasing cycle number (Fig. S9b†). Additionally, the capacity of stored electrons also reduced from 0.6033 mmol g⁻¹ to 0.5218 mmol g⁻¹. The above results suggested that 80% of the performance of h-WO₃ \cdot 0.46H₂O was retained after 10 cycles, reflecting the good recyclability of the catalyst. Furthermore, the cycle experiments had no discernible effects on the crystal phase of h-WO₃ \cdot 0.46H₂O (Fig. S10†). The tremendous potential of h-WO₃ \cdot 0.46H₂O in practical applications is indicated by its exceptional performance and structure stability.

After the analysis and discussion mentioned above, the reasonable cycle mechanism for h-WO₃ \cdot 0.46H₂O under all-weather conditions is proposed in Fig. 6h. Under the excitation of visible or solar light, the electron storage effect initiated by the process of W(vi) \rightarrow W(v) increases the local electronic



Fig. 8 The degradation pathway of DNBP in the illuminated h-WO₃·0.46H₂O/H₂O₂ system in the dark and the toxicity evolution of intermediates in the degradation pathways estimated via ECOSAR.

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density of h-WO₃·0.46H₂O and the protons originating from lattice water located in the dodecagon tunnels along the z axis interact with the nearby bridging oxygen atoms in the lattice, which causes the colour of h-WO3 · 0.46H2O to change from white to blue and the distortion of its lattice in the horizontal and vertical directions, respectively. The lattice change also promotes the entry of water molecules from outside into the lattice of h-WO₃·0.46H₂O to replenish the lattice water molecules consumed by the photochromic effect. In the dark, the exogenous H2O2 molecule tends to adsorb horizontally on the surface of illuminated h-WO₃·0.46H₂O under thermodynamic and kinetic driving forces. Subsequently, the adsorbed H₂O₂ molecule is activated by the stored electrons in illuminated h- $WO_3 \cdot 0.46H_2O$ and O_2^- , OH and O_2^- are efficiently produced. Accordingly, it is possible to see the excellent DFR activity of illuminated h-WO₃ \cdot 0.46H₂O. In the bargain, the altered colour and structure of h-WO₃·0.46H₂O due to illumination is also restored by the depletion of the stored electrons.

The transformation of the structure of organic pollutant molecules during degradation is closely related to the reactivity of its different moieties. As shown in Fig. 7a, the DNBP molecule was constructed and optimized by GaussView before calculation (the details of the calculation are shown in Text S1.3[†]). After that, a series of theoretical calculations based on density functional theory (DFT) were carried out to investigate the reactivity of different regions of the DNBP molecule. The C (1-6), N (12), O (11), O (14) and O (15) atoms in the DNBP molecule have the majority of the HOMO and LUMO orbitals (Fig. 7b and c), demonstrating that these atoms are more susceptible to assault.51-53 In addition, the electrostatic surface potential (ESP) distribution diagram (Fig. 7d) of DNBP intuitively showed the location with red color of negatively charged centers in the molecule.54 The two nitro groups are directly attached to the benzene ring and have a higher surface electron cloud density, thus making them susceptible to attack by species with strong oxidizing capacity.55,56 According to the Fukui indices of the DNBP molecule shown in Fig. 7e, the C (1), C (2), C (4-7), N (12 and 13), O (14) and O (16) atoms have a high Fukui (0) index, which indicate that these sites have the theoretically higher radical reactivity.57-59 The C (4), C (6, 7), N (12 and 13) and O (11, 14-17) have considerable electrophilic reactivity as indicated by their Fukui (-) indexes.58-61

The genuine degradation paths of DNBP molecules in the illuminated h-WO₃·0.46H₂O/H₂O₂ system are obtained by GC-MS tests in addition to DFT calculations. Based on a rigorous analysis of the MS spectra (Fig. S11 and Table S3[†]) of the reaction intermediates, two main rational pathways are proposed (Fig. 8). As displayed in Fig. 8, N (12) is assaulted firstly by the ROS and the o-nitro group leaves. Then, the obtained 2-*sec*-4-nitro-butylphenol intermediate undergoes two routes of degradation reactions simultaneously. On the one hand, the N (13) site is next attacked to cause the departure of the p-nitro group (path 1), and the formed intermediate is immediately followed by a series of oxidation reactions (the reaction sites are C (5–7)) and is finally mineralized; on the other hand, the C (7) atom as the reactive site is oxidized (path 2) and then continuous oxidation and ring opening with N (13)

and C (1-4, 6) atoms as the reaction centers occur one after another. The pathway 3 seems to be much easier than paths 1 and 2: the C (7) site is oxidized firstly by ROS and the formaldehyde group is formed. Next, the C (7) site stays as the reaction site as the intermediate formed in the previous step passes through oxidation and hydroxylation reaction. Afterwards, C (5) and C (6) as reactive sites are attacked and the ring opening reaction is induced. Finally, the product generated by ring opening reaction is mineralized. It is obvious that the reactive sites obtained from the results of DFT calculation can basically match with the above degradation pathways. In the bargain, an intuitive conclusion can also be seen in Fig. 8 that the toxicity of intermediates in the degradation process of DNBP gradually decreases with the increase of reaction time, which indicated that the removal path of DNBP by the DFR process was efficient and safe.

4 Conclusions

In conclusion, a hexagonal tungsten oxide hydrate (h-WO3·0.46H2O) was synthesized and intensively investigated based on a series of characterizations and experiments. By the Rietveld refinement of the XRD spectra of $h-WO_3 \cdot 0.46H_2O$, the lattice water molecules can be presumed to be in the dodecagon tunnels along the z axis and reside between W-O layers along the z axis. Under visible light illumination, h-WO₃ \cdot 0.46H₂O showed slow discoloration (white-blue) and the lattice water gradually decomposed. Owing to the strong interaction between the protons originating from lattice water and the bridging oxygen atoms (W-O-W), the blue colour of illuminated h- $WO_3 \cdot 0.46H_2O$ can remain stable for a long time (>24 h). More interestingly, the stored electrons due to photochromic effect in illuminated h-WO₃·0.46H₂O can effectively activate H₂O₂ and quickly degrade DNBP in the dark. On the basis of the experimental and theoretical findings, the mechanism of ROS (O_2^- , 'OH and ¹O₂) formation and potential pathways for DNBP degradation were eventually clarified. Our proposed mechanism, which is more reasonable and excludes the 1e⁻/2e⁻ ORR pathway in the dark, is different from the more nebulous allweather processes of WO₃-containing photocatalysts that have been reported. Our work demonstrates that the photo-storage electrons in h-WO₃·0.46H₂O activate H₂O₂ to form ROS and excludes the possibility that it generates reactive oxygen species through the mechanism previously reported in the literature, which may attract more researchers to pay attention to the reaction ability of photo-storage electrons in WO₃. In addition, we have only studied the activation effect of photo-storage electrons in h-WO₃ \cdot 0.46H₂O on H₂O₂, but more valuable chemical processes may also be triggered by the photo-storage electrons, which may have more scientific questions to be explored.

Author contributions

Hao Huang: conceptualization, data curation, methodology, validation, formal analysis, riting – original draft preparation, investigation, visualization, writing – review & editing,

resources. Hui-Long Wang: conceptualization, supervision, project administration, resources, writing – review & editing, funding acquisition. Wen-Feng Jiang: writing-review & editing, supervision, project administration.

Conflicts of interest

There are no conflicts to declare.

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