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Hyper oxygen incorporation in CeF₃: a new intermediate-band photocatalyst for antibiotic degradation under visible/NIR light†

Bing Han,^a Siqu Yu,^{ID a} Dian Zhao,^{ID ab} Yunchao Lou,^a Jiayang Gao,^a Zhe Liu,^a Zhiyu Wang^{ID *a} and Guodong Qian^{ID *a}

Intermediate-band semiconductors perform functions similar to natural photosynthesis by combining two photons to achieve a higher electron excitation. In this study, a strategy was developed to prepare a high oxygen-doped CeF₃ (CeF₃-O) nanomaterial that exhibits photocatalytic activity under visible/NIR light for the first time. The homogeneous doping oxygen atoms were verified to efficiently modify the band structure of CeF₃. DFT calculation predicted the formation of an intermediate band within CeF₃ upon homogeneous doping of O at interstitial sites. The interaction between F and O atoms generates an intermediate band, which divides the total bandgap of CeF₃-O into two sub-bandgaps at about 1.7 eV and 2.9 eV, enabling CeF₃-O photocatalysis under visible light and NIR light. Reflectance spectra evidenced that the same bandgaps exist. The photocatalytic activities of CeF₃-O were tested by wavelength-controlled light. The rate constants of TC-HCl photodegrading under visible/NIR light are $12.85 \times 10^{-3} \text{ min}^{-1}$ and $1.28 \times 10^{-3} \text{ min}^{-1}$, respectively. The two-step electron transfer was also obviously confirmed in visible-light photocatalysis. In conclusion, the high oxygen doping builds a more applicable band structure of CeF₃-O for photocatalytic performance, charge transfer and special light response for visible/NIR light.

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1. Introduction

A large number of semiconductor photocatalysts have been investigated to achieve the goal of degrading organic pollutants into non-toxic products by solar energy.^{1–3} On the basis of photocatalysis, the photodegradation efficiency of semiconductor-based photocatalysts is mainly determined by its bandgap, the position of conduction band (CB) and valence band (VB), and photogenerated carrier transfer.^{4,5} Due to poor penetration of ultraviolet light in water, long-wave sunlight is more suitable for organic pollutant photodegradation. Meanwhile, for improving the solar utilization, semiconductor photocatalysts have been modified by the methods of ion doping, Z-scheme and quantum dots to broaden the solar response range.^{6–9}

The visible light activity of semiconductor photocatalysts, in particular TiO₂, has been improved by metal and non-metal doping for several decades.^{10,11} The doping energy levels in

this band structure just try to decrease the bandgap in order to extend spectral absorption to the visible region. However, the potential of the [•]OH/H₂O reaction is 2.72 eV vs. NHE. This demands that semiconductor photocatalysts take on a high enough VB, where holes react with water molecules to generate oxygen radicals. The reduced bandgap of the ion-doped photocatalyst is unconvincing to the generation of oxygen radicals. Thus, the Z-scheme heterojunction has been rapidly developed for the photodegradation of organic pollutants to combine two low-energy photons to achieve a higher energy electron excitation with a broader sunlight illumination.^{12,13}

With the same two-photon absorbing manner, the intermediate band scheme has increased the current of solar cells, which implements the two-photon working principle.¹⁴ Different from the Z-scheme, intermediate band scheme does not spatially separate the two absorptions, so that a smaller electron-transfer barrier exists, taking advantage of full electronic transferring.¹⁵ An important way to achieve the intermediate band scheme is to introduce intermediate band levels in a host semiconductor via hyper doping.¹⁶ Different from the light ion doping of the previously reported photocatalyst TiO₂, this essentially requires a high enough concentration of doping ions to ensure that the inserted levels could have a delocalized character to form a true band. However, it has been verified that these doping ions can reduce nonradiative recombination.¹⁷ In

^aState Key Laboratory of Silicon Materials, Department of Materials Science and Engineering, Zhejiang University, Hangzhou 310027, China. E-mail: wangzhiyu@zju.edu.cn; gdqian@zju.edu.cn

^bKey Laboratory of the Ministry of Education for Advanced Catalysis Materials, College of Chemistry and Life Sciences, Zhejiang Normal University, Jinhua, 321004, China

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the past years, several intermediate band scheme materials are used as solar cells and photocatalysts.

The rare-earth compounds with huge characteristic f-f and f-d transitions are perfect luminescent materials with high photoelectric conversion efficiency and long lifetime of excited state due to its rich electrons and energy levels.^{18,19} All the time, the research of rare earth photocatalysts focus on CeO₂ and rare earth-based perovskites, some in connection with CeF₃.^{20–22} CeF₃-based composite photocatalysts could convert visible light into ultraviolet light by efficient upconversion emissions of CeF₃, enhancing the photocatalytic activity under visible light.^{23,24} It has been hardly regarded as a feasible photocatalyst by itself and rarely used in NIR light photocatalysis due to its wide bandgap.²⁵ The larger lattice space of CeF₃ makes it to be an ideal host semiconductor material for hyper doping. The mechanism of the intermediate band scheme shows a viable way to synthesize a novel CeF₃ photocatalyst. Considering about the interaction between Ce atoms and other non-metal atoms, oxygen may be easier to be introduced as a doping atom and may easily modify CeF₃.

Modified CeF₃ applied in photocatalysis is an important extension for photocatalysts. Herein, we focus on a special process to investigate a CeF₃-O photocatalyst (CeF₃ with a hyper oxygen doping). Open-framework H_{25.5}(NH₄)_{10.5}Ce₉O₂₇F₁₈ was required as the precursor and synthesized under simpler conditions than that ever reported.²⁶ The introduction of oxygen efficiently modified the electronic band gap structure, generating an intermediate band. An obvious two-step electron transfer was observed in the photocatalysis process. CeF₃-O with high-concentration doped oxygen up to 18.7 mol% exhibited a porous structure and activity for tetracycline hydrochloride (TC-HCl) photodegradation under both visible and NIR light.

2. Methods

2.1 Materials and analysis methods

Cerium(IV) diammonium nitrate ((NH₄)₂Ce(NO₃)₆), ammonium fluoride (NH₄F), and tetracycline hydrochloride (TC-HCl) were purchased from Sinopharm Chemical Reagent Co. Ltd. All reagents were of analytical grade and used without further purification. Deionized water was used throughout the experiments.

SEM (scanning electron microscopy) images were acquired using a Hitachi S4800 field-emission scanning electron microscope. XRD (X-ray diffraction) patterns were recorded using a Shimadzu XRD-7000 X-ray diffractometer equipped with a Cu Kr radiation source at a scanning rate of 1° min^{−1}. PL (photoluminescence) spectra were recorded using a Hitachi F-4600 fluorescence spectrometer. UV-vis spectra and UV-vis diffuse reflectance spectra were recorded using an Agilent Cary 5000 spectrophotometer. XPS (X-ray photoelectron spectroscopy) spectra were recorded using a Thermo Fisher Scientific ESCA-LAB 250XI and the binding energies were calibrated by C 1s 284.8 eV. The BET surface area was estimated using a Micromeritics ASAP 2020 adsorption porosimeter based on the nitrogen isotherms at 77 K. TG (thermogravimetric) analysis

was carried out using a NETZSCH TG 209 thermal analyzer at a heat rate of 5 °C min^{−1} in the air. Photoelectrochemical estimates were completed using a Chenhua CHI660E electrochemical workstation.

2.2 Synthesis of CeF₃-O nanoparticles

In a typical synthesis procedure, 0.27 g (NH₄)₂Ce(NO₃)₆ was dissolved in 20 mL deionized water. Subsequently, 0.445 g NH₄F was added under rapid stirring and white precipitates were formed immediately. The mixture was transferred into a Teflon bottle and held in a stainless-steel vessel to further seal. Then, the stainless-steel vessel was placed in a thermostatted oven and heated at 150 °C for 6 h to produce the precursor H_{25.5}-(NH₄)_{10.5}Ce₉O₂₇F₁₈. After heating, the precursor was collected, washed with deionized water and dried. Finally, the dried precursor was calcined at 260 °C, 270 °C and 280 °C, respectively for 1 h at a heat rate of 1 °C min^{−1} in the air atmosphere.

2.3 Photocatalytic degradation and active species measurements

The photocatalytic degradation was assessed in a 50 mL TC-HCl (20 mg L^{−1}) solution with 20 mg photocatalyst. The reactant mixture was cooperated with magnetic stirring and kept in darkness for 30 min to achieve the adsorption-desorption equilibrium between CeF₃-O and TC-HCl. Then, the solution was illuminated using a PLS-SXE 300 Xenon lamp with a UV-cutoff filter (>400 nm). After different irradiation intervals, 2 mL sample was taken and followed by centrifugation. The TC-HCl solution concentration was analysed by UV-vis spectrophotometry. The RhB photodegradations followed the same method.

2.4 Photoelectrochemical test

Photoelectrochemical measurements were performed with a standard three-electrode configuration using CeF₃-O-coated ITO (area: 1 cm²) as the working electrode, a Pt plate as the counter electrode, a saturated calomel electrode as the reference electrode, and 0.1 M Na₂SO₄ as the electrolyte. A xenon lamp was utilized to supply light, which was cut off by filters ($\lambda > 400$ nm or $\lambda > 680$ nm). First, 10 mg photocatalyst was dispersed by 1800 μ L ethanol and 100 μ L Nafion solution (5 wt%) under sonication to make the homogeneous ink, then 50 μ L ink was pipetted on the ITO surface to gain films.

2.5 Theoretical calculations

Density functional theory (DFT) calculations were performed using the Vienna *ab initio* simulation package (VASP) by generalized gradient approximation (GGA) with Perdew–Burke–Ernzerhof (PBE) functions.^{27–30} Spin polarization was considered and the energy cutoff was set to 520 eV. The energy and force convergence criterion were set to 1×10^{-7} eV and 1×10^{-4} eV Å^{−1}, respectively. The DFT+U method proposed by Dudarev *et al.* was adopted with a (*U*–*J*) value of 5.0 eV to describe the strongly correlated f-electrons of Ce.³¹ van der



Waals corrections (DFT-D2 method proposed by Grimme) were also incorporated to characterize the long-range interactions.³²

3. Results and discussion

3.1 Characterization of the obtained CeF₃-O

We prepared the H_{25.5}(NH₄)_{10.5}Ce₉O₂₇F₁₈ precursor by mixing and heating (NH₄)₂Ce(NO₃)₆ and NH₄F aqueous solution (Fig. S1a†). Then, the dried precursor was calcined at various temperatures. The detailed XRD patterns of the obtained samples are shown in Fig. S1b†. The diffraction peaks of samples match well with CeF₃. However, after calcination at 260 °C, the precursor decomposes incompletely; while after calcination at 280 °C, CeO₂ is observed. Fig. 1a shows the characteristic diffraction peaks of CeF₃ obtained by calcination at 270 °C. Interestingly, the positions of diffraction planes in this CeF₃ are slightly different from the standard (Table 1), which indicates that the lattice distortions are formed.

To evaluate the lattice distortion of this CeF₃, XPS spectra of elemental O, F, N and Ce were provided. The doping ratio of O is up to 18.7 mol% (Table S1†), which may have a huge impact on the lattice. Two typical peaks of O at 529.0 and 531.6 eV (labelled as O1, O2) in Fig. 1b are attributed to lattice oxygen and chemisorbed oxygen, respectively.³³ The Ce 3d XPS spectra (Fig. S2†) display Ce 3d_{3/2} and 3d_{5/2} states assigned to more Ce³⁺ and fewer Ce⁴⁺.^{34,35} The main peak of F 1s (Fig. 1c) at 684.4 eV shifts towards the lower binding energy compared with the standard value of 685.7 eV, which suggests the banding between Ce and the doping O. Moreover, no crystal phase of rare earth oxides is surveyed in the XRD patterns, showing that doped O

was homogeneously distributed at interstitial sites. Thus, this CeF₃ doped with O is named by CeF₃-O. The CeF₃-O appears in pale pink (Fig. 1a inset) to enhance the absorption of visible light. The N 1s peak does not appear in the XPS spectrum (Fig. 1d), suggesting that the colour's diversity is due to the appearance of oxygen. The FTIR spectra (Fig. S3†) further confirm the structural change. Compared with the commercial CeF₃, two sharp bands of CeF₃-O located at 3280 and 1426 cm⁻¹ with strong intensity are attributed to the stretching and bending vibrations of -OH, respectively. It forms from the oxygen of CeF₃-O, which have affinities with water molecules in the air. The typical peak at 574 cm⁻¹ is the Ce-O stretching vibration. The above-mentioned results reveal that the oxygen atom is the chief reason of lattice distortion, and its hyper doping ratio and Ce-O bonding may create the intermediate band and have great influence on the performance of the photocatalyst.

The precursor H_{25.5}(NH₄)_{10.5}Ce₉O₂₇F₁₈ has a special open-framework structure, which is constituted by six CeO₆F₂ dodecahedra (Fig. 2a and S4a†).²⁶ The basic building unit of CeO₆F₂ provides a major factor for homogeneous oxygen-doped CeF₃-O preparation. Further characterization by TEM obviously reveals that the precursor presents a hollow rhombohedral structure at a scale ranging from 100–400 nm (Fig. 2b and S4b†). During the calcination process, the precursor gradually decomposed into smaller nanoparticles. The rhombohedral structure is still kept and built up by CeF₃-O nanoparticles, when calcination was completed (Fig. 2d). Besides, the element mappings demonstrate the homogeneous distribution of cerium, fluorine and oxygen elements in CeF₃-O (Fig. 2e–g). It was also found that the surface area of CeF₃-O was 39.3 m² g⁻¹, and a type-IV isotherm with an H3 hysteresis loop is presented in Fig. 2c. Such porous structures with a larger surface area might play a positive role in photocatalysis, which can possess efficient exposure and active sites with the substrate.

3.2 Energy gap analysis and intermediate band

The first-principles pseudopotential methods based on DFT were carried out to understand the influence of doping O on the band structure and electronic properties of CeF₃-O. On the basis of the XPS test, it is assumed that there is an oxygen atom in the CeF₃-O structure doping per CeF₃ at interstitial sites. The model is shown in Fig. 3a and S12.† The calculated band structure and DOS of pristine CeF₃ and CeF₃-O (Fig. 3 and S5†) confirm the changes attributed to oxygen doping. Unlike the direct band gap of CeF₃, CB and VB are located at different Brillouin zones. CeF₃-O changes into an indirect band gap. A new energy level generates at about 1.2 eV as an intermediate band. The density of states (DOS) diagrams of CeF₃-O (Fig. 3d and S7†) are displayed to reveal the contributions of every participant orbital to

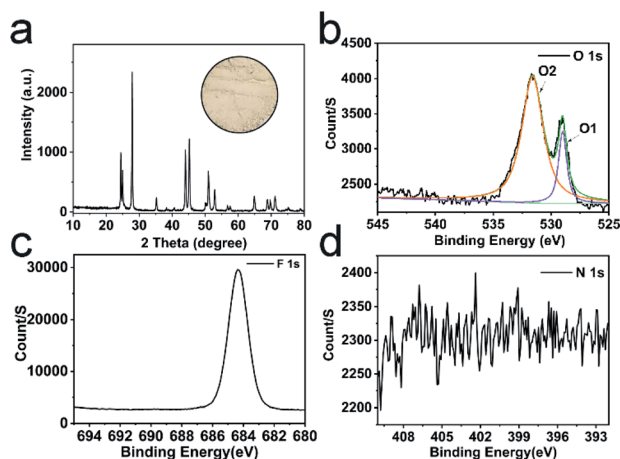


Fig. 1 XRD pattern of CeF₃ calcined at 270 °C (JCPDS card no. 08-0045), the inset is the optical image of CeF₃ calcined at 270 °C (a); XPS spectra of O 1s (b), F 1s (c) and N 1s (d).

Table 1 Difference between CeF₃ for 270 °C calcination and standard

Lattice plane	(110)	(300)	(302)	(221)	(214)	(304)	(115)	(411)
2θ (degree) of standard	25.06°	44.05°	51.02°	52.96°	64.98°	68.90°	69.74°	71.30°
2θ' (degree) of 270 °C	25.00°	43.98°	50.92°	52.88°	64.88°	68.86°	69.68°	71.14°



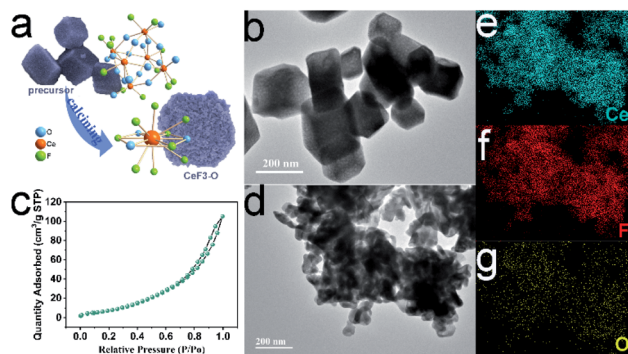


Fig. 2 Schematic of the structural modification from the precursor $\text{H}_{25.5}(\text{NH}_4)_{10.5}\text{Ce}_9\text{O}_{27}\text{F}_{18}$ to $\text{CeF}_3\text{-O}$ (a); TEM of the precursor $\text{H}_{25.5}(\text{NH}_4)_{10.5}\text{Ce}_9\text{O}_{27}\text{F}_{18}$ (b) and $\text{CeF}_3\text{-O}$ (d), N_2 adsorption/desorption isotherm of $\text{CeF}_3\text{-O}$ (c); elemental mapping of cerium, fluorine and oxygen in $\text{CeF}_3\text{-O}$ nanoparticles (e–g).

the band. It can be seen that the intermediate band mainly comprises the dopant O 2p and F 2p unoccupied molecular orbitals by reason of the close-range between O_2^- and F^- ions. The maximum of VB is composed of dopants O 2p, F 2p and Ce 4f occupied molecular orbitals. The minimum of CB principally originates from the Ce 5d and 4f states. The charge density distributions of pristine CeF_3 and $\text{CeF}_3\text{-O}$ on the [001] plane agree with the previous calculations of bandstructure and DOS (Fig. S8†). For pristine CeF_3 , electronic charges transfer from electropositive Ce atoms to the adjacent electronegative F atoms, forming ionic bonds. For $\text{CeF}_3\text{-O}$, O exhibits lighter electronegativity and more electrons transfer from Ce atoms to other types of atoms.

Moreover, two new energy gaps exist. The broader one between the intermediate band and VB is 2.9 eV, which is the main reason for the pale pink colour. The narrow one calculated by the intermediate band and CB is 1.8 eV. Both of them enable $\text{CeF}_3\text{-O}$ with visible light absorption. Simultaneously, when O is doped into CeF_3 , the electrons transferring from the VB to CB can be effortlessly to jump a tiny energy gap. That is favourable for the electron–hole pair generation and separation. These calculation results may indicate that $\text{CeF}_3\text{-O}$ possesses photocatalytic activity in the visible-light region.

These calculated values based on DFT are generally verified. Fig. 3b shows the optical bandgaps of $\text{CeF}_3\text{-O}$ generated from the diffuse reflectance spectrum by the Kubelka–Munk equation.^{36,37} The chief bandgap estimated from the intercept of the tangent to the plot is around 2.9 eV, while a tail bandgap was evaluated to be 1.7 eV, making the possibility of TC-HCl photodegradation under NIR light irradiation. The bandgap values of reflectance spectra are well coincided with the values obtained from the calculation.

3.3 Photocatalytic performance of $\text{CeF}_3\text{-O}$ under visible/near-infrared light and its two-photon absorption

The PL emission of $\text{CeF}_3\text{-O}$ is completely quenched compared to commodity CeF_3 (Fig. 4a). The commodity CeF_3 exhibits a heavy emission peak at 330 nm ascribed to the high electron–hole pair

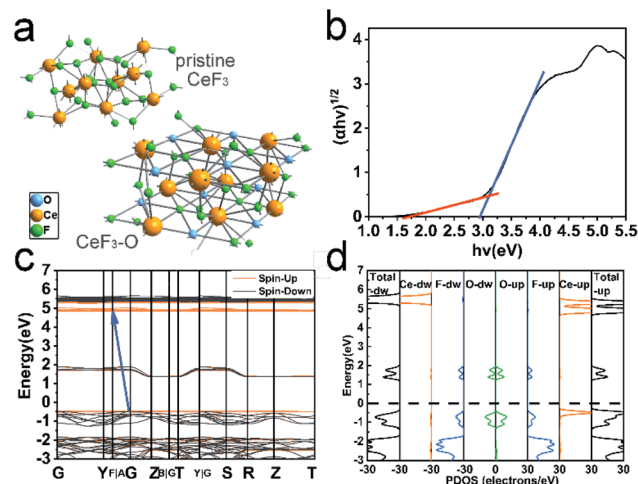


Fig. 3 Schematic of pristine CeF_3 and $\text{CeF}_3\text{-O}$ crystal structures. The interacting O atoms were labeled in blue (a); the optical band gaps of $\text{CeF}_3\text{-O}$ using the Kubelka–Munk function (b); DFT-calculated band structure of $\text{CeF}_3\text{-O}$, details in Fig. S5† (the blue arrow points from VB to CB) (c); the total and partial DOS of $\text{CeF}_3\text{-O}$ (d).

recombination rate, while $\text{CeF}_3\text{-O}$ shows a very low intensity. In the absence of O_2^- ions as nonradiative centres, electrons and holes remain spatially diffusing. This results in increasing the life time of charge carriers and demonstrates the appearance of photocatalytic performance by $\text{CeF}_3\text{-O}$.

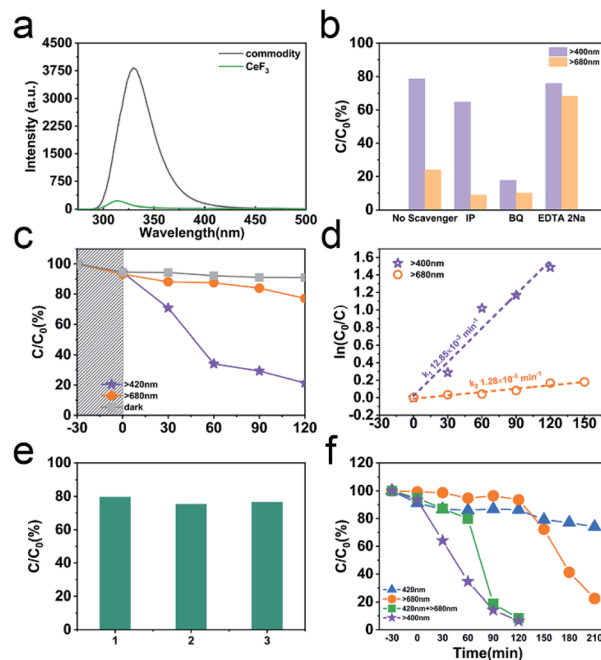


Fig. 4 PL spectra of commodity CeF_3 and $\text{CeF}_3\text{-O}$ with an excitation wavelength of 251 nm (a); free-radical trapping experiments of $\text{CeF}_3\text{-O}$ under different cutoff filters (darkness, $\lambda > 400$ nm and $\lambda > 680$ nm) (b); photodegradation of TC-HCl by $\text{CeF}_3\text{-O}$ under different cutoff filters (darkness, $\lambda > 400$ nm and $\lambda > 680$ nm) (c); photocatalytic kinetic profile (d); reusability of $\text{CeF}_3\text{-O}$ to degrade TC-HCl under visible light (e); photodegradation of RhB by $\text{CeF}_3\text{-O}$ under different conditions ($\lambda > 400$ nm, $\lambda = 420$ nm and $\lambda > 680$ nm) (f).



For evaluating the photocatalytic performance, a significant TC-HCl photocatalytic degradation efficiency is observed in CeF₃-O (Fig. 4c). In the darkness, the TC-HCl absorption of CeF₃-O is about 9.0%, while the process of CeF₃-O degrading TC-HCl takes 120 min and boasts a 78.7% efficiency rate under visible light. Under the same conditions, the degrading ratio under NIR light is 22.8%. The degradation with time is fitted by a pseudo-first-order kinetic model:³⁸

$$\ln\left(\frac{C_0}{C}\right) = kt,$$

where k (min⁻¹) is the apparent reaction rate constant of degradation; C_0 (mg L⁻¹) and C (mg L⁻¹) are the initial concentration of TC-HCl and the concentration of TC-HCl at time, respectively. The apparent reaction rate constants k are shown in Fig. 4d. The coefficients of regression (R_2) are not higher than 0.94 due to the poor fitting of all the experimental data. The rate constant of CeF₃-O under visible light is 12.85×10^{-3} min⁻¹, which is about ~10 times greater than that under NIR light. The degradation efficiency remained at 76.5% after three cycles under visible light, suggesting the stability of CeF₃-O (Fig. 4e). As a contrast, commercial CeF₃ scarcely has photocatalysis activity both under visible and NIR light (Fig. S10†). The photodegradation of TC-HCl by CeO₂ is slightly higher than commercial CeF₃. As a result, it confirms the visible-light-response photocatalytic properties of CeF₃-O.

To further understand the photocatalytic mechanism, the trapping experiments in the photocatalytic process under visible and NIR light are displayed in Fig. 4b. BQ, EDTA 2Na and IP are used to scavenge $\cdot\text{O}_2^-$, h^+ and $\cdot\text{OH}$ species, respectively.^{39–41} The addition of IP scavenges $\cdot\text{OH}$ and has little effect on the TC-HCl degradation. The $\cdot\text{OH}$ contribution is exiguous compared to $\cdot\text{O}_2^-$. By adding EDTA 2Na, to a more extent, the reaction is 75.9% of TC-HCl degradation. The degradation in the presence of BQ is 17.6%, suggesting that the photocatalytic performance is suppressed significantly and $\cdot\text{O}_2^-$ is the main active species in this photocatalytic process. Like the results under visible light, the degradations under NIR light in the presence of IP and BQ are both below 10%, showing that the photocatalytic performance is almost terminated. It testifies that there are barely $\cdot\text{OH}$ and $\cdot\text{O}_2^-$ at this process. The addition of EDTA 2Na captures h^+ and reduces the carrier recombination, increasing the degradation up to 68.2%. No matter under visible or NIR light, $\cdot\text{O}_2^-$ is the key to degrade TC-HCl. Thus, photogenerated electron in a suitable potential of CB can engender $\cdot\text{O}_2^-$ to participate in the redox reaction for TC-HCl degradation.

To further figure out the intermediate band, other degradations of RhB are tested by using different cutoff filters (Fig. 4f). The degradations are just 25.9% ($\lambda = 420$ nm) and 77.6% ($\lambda > 680$ nm) after illumination lasting 240 min, individually. By the dual light sources, the degradation is completed in 120 min, which is concomitant with the results in the range of visible light. This means that the photocatalysis follows obvious two-photon absorption. When only the light ($\lambda > 680$ nm) irradiates, electrons absorb 1.7 eV energy and transfer from VB to intermediate band, bringing out an oxidation–reduction reaction. However, this

progress is of low efficiency. After addition of the irradiation at 420 nm, the rate of photocatalytic reaction rapidly increases, which may attribute to the second photon absorption. In this process, redundant electrons absorb 2.9 eV energy and transfer from the intermediate band to CB, carrying on another oxidation–reduction reaction. The intermediate band scheme makes the photocatalytic reaction more efficient under visible light.

3.4 Electrochemistry properties and the proposed photocatalytic mechanism

Faster separation and transfer of photogenerated carriers can achieve much better photocatalysis property. The Nyquist impedance plots for CeF₃-O under light irradiation are displayed (Fig. 5a). The arc radius of CeF₃-O under visible light is smaller, which demonstrates that CeF₃-O possesses a finer electronic conductivity in photoexcited state and a less obstruction for the transfer of photogenerated carries.^{42,43} A Mott–Schottky test was applied to describe the carrier density upon the Schottky barrier between electrolytes and electrodes, which could further confirm semiconductor types and flat band potential. Fig. 5b exhibits the Mott–Schottky plots for CeF₃-O, which displays a positive slope. It is the characterization of the n-type semiconductor. Based on the following equation:⁴⁴

$$\frac{1}{C^2} = \frac{2}{q\epsilon\epsilon_0 N_d} \left(E - E_{fb} - \frac{K_b T}{q} \right)$$

The flat band potential of CeF₃-O can be calculated to be -1.00 eV vs. SCE, which is -0.76 eV vs. NHE. CB can be negative about 0.1–0.4 eV in comparison to flat band.^{45,46} Thus, CB of the CeF₃-O is noted at -0.86 eV. According to the band structure and active species experiments, the VB position of CeF₃-O ($+3.74$ eV vs. NHE) is more positive than $E(\cdot\text{OH}/\text{OH}^-)$ ($+2.38$ eV vs. NHE) and $E(\cdot\text{OH}/\text{H}_2\text{O})$ ($+2.72$ eV vs. NHE). Holes generated in VB can easily react with water molecules adsorbed on the surface of photocatalyst or hydroxyls to form $\cdot\text{OH}$ radicals. Meanwhile, the CB position (-0.86 eV) is negative by contrast with $E(\text{O}_2/\cdot\text{O}_2^-)$ (-0.33 eV vs. NHE); electrons in CB react with O_2 to engender $\cdot\text{O}_2^-$ radicals. For the intermediate band, the position $+2.04$ eV is above $E(\text{H}^+, \text{O}_2/\text{H}_2\text{O}_2)$ ($+0.94$ eV vs. NHE), and electrons could react with H^+ and O_2 to produce H_2O_2 , slowly degrading TC-HCl.

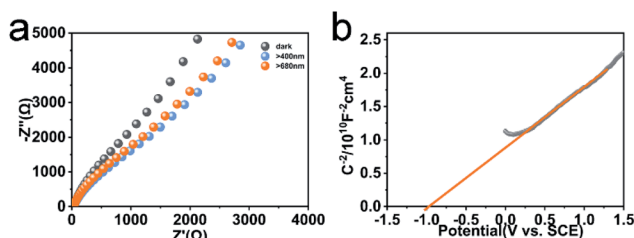


Fig. 5 EIS spectra of CeF₃-O without or with light ($\lambda > 400$ nm and $\lambda > 680$ nm) (a). Mott–Schottky plots of the CeF₃-O (b).



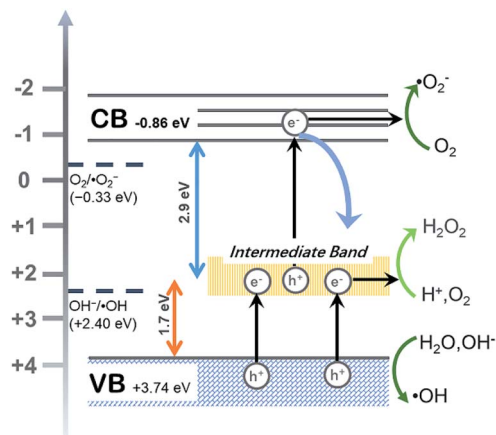


Fig. 6 Proposed photocatalytic mechanism by the $\text{CeF}_3\text{-O}$ photocatalyst under visible light irradiation.

All above decidedly elucidate the reason for the visible-light-spectrum-response photocatalytic activity of $\text{CeF}_3\text{-O}$. The whole process can be broken into three steps (illustrated in Fig. 6):

(i) The intermediate band is generated in the new band structure, which not only can be an electron acceptor but also an electron donor. (ii) Electrons absorb a certain energy and transfer from VB to the intermediate band, leaving an electron vacancy; the electron-hole pair separates. Parts of oxidation-reduction reaction run in VB and a new CB of intermediate band. (iii) The redundant electrons absorb another certain energy and transfer once more from the intermediate band to CB to occupy the electron vacancy, producing the electron hole pair in the CB and VB. The electron-hole pair separates again, resulting in another oxidation-reduction reaction.

4. Conclusion

In summary, $\text{CeF}_3\text{-O}$ is presented for the first time as a photocatalyst under visible and NIR irradiation. The special process to investigate $\text{CeF}_3\text{-O}$ is based on the way of thermal control decomposing the precursor $\text{H}_{25.5}(\text{NH}_4)_{10.5}\text{Ce}_9\text{O}_{27}\text{F}_{18}$, which is synthesized differently from that ever reported. $\text{CeF}_3\text{-O}$, with high-concentration oxygen dopants up to 18.7 mol%, exhibits a porous structure and favourable activities for TC-HCl photodegradation. Besides, the test confirms that the main active species in visible light photocatalysis is $\cdot\text{O}_2^-$ and NIR light is e^- .

The introduction of oxygen is confirmed to largely modify the electronic band gap structure, which is verified by simulated calculation and experimental results. Compared to CeF_3 , $\text{CeF}_3\text{-O}$ comprises a new intermediate band. Two narrow band gaps are formatted at about 1.7 eV and 2.9 eV, making photocatalysis follow the obvious two-step electron transfer. More importantly, these results indicated that oxygen doping can modify the band structure and electronic property of $\text{CeF}_3\text{-O}$ for photocatalytic performance, transfer charge easily as well as broaden the light response region, even in the near-infrared region. It is believed that this method has great significance for exploring an enlightening pathway to design and synthesize porous photocatalysts in the visible and NIR regions.

Conflicts of interest

There are no conflicts to declare.

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