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Photocatalytic degradation of methyl orange by CeO2 and Fe-doped CeO2 films under visible light irradiation

Abstract

Undoped CeO2 and 0.50-5.00 mol% Fe-doped CeO2 nanoparticles were prepared by a homogeneous precipitation combined with homogeneous/impreganation method, and applied as photocatalyst films prepared by a doctor blade technique. The superior photocatalytic performances of the Fe-doped CeO2 films, compared with undoped CeO2 films, was ascribed mainly to a decrease in band gap energy and an increase in specific surface area of the material. The presence of Fe3+ as found from XPS analysis, may act as electron acceptor and/or hole donor, facilitating longer lived charge carrier separation in Fe-doped CeO2 films as confirmed by photoluminescence spectroscopy. The 1.50 mol% Fe-doped CeO2 film was found to be the optimal iron doping concentration for MO degradation in this study.

Keywords

light, photocatalytic, degradation, methyl, orange, irradiation, ceo2, visible, fe, doped, films, under

Disciplines

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Photocatalytic Degradation of Methyl Orange by CeO₂ and Fe–doped CeO₂ Films under Visible Light Irradiation

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Undoped CeO_2 and 0.50-5.00 mol% Fe-doped CeO_2 nanoparticles were prepared by a homogeneous precipitation combined with homogeneous/impreganation method, and applied as photocatalyst films prepared by a doctor blade technique. The superior photocatalytic performances of the Fe-doped CeO_2 films, compared with undoped CeO_2 films, was ascribed mainly to a decrease in band gap energy and an increase in specific surface area of the material. The presence of Fe^{3+} as found from XPS analysis, may act as electron acceptor and/or hole donor, facilitating longer lived charge carrier separation in Fe-doped CeO_2 films as confirmed by photoluminescence spectroscopy. The 1.50 mol% Fe-doped CeO_2 film was found to be the optimal iron doping concentration for MO degradation in this study.

rganic pollutants from the textile industry represent a major source of water contamination. About 15% of commercially used dyes are released without treatment¹, while those that are required either a chemical or energy intensive processing which leads to further strain on the environment. An inherent requirement of most of these dyes is that they are stable for long peroids of time in their application. Unfortunately, this also makes degradation (particularly biodegradation) of their excess difficult. Methyl orange (MO) is a simpleazo dye. Azo dyes are widely used in textile, printing, pharmaceutical, and research laboratories. It can enter the human body through skin, and can cause the death of lung tissues, fast heart rate, and vomiting². For these reasons, MO was chosen as a model dye for photocatalytic degradation in this research.

Advanced oxidation processes based on semiconductor oxide photocatalysts are an effective method for dye degradation³, with various oxide semiconductors such as TiO_2 , ZnO, and WO_3 being used in heterogeneous photocatalysis⁴⁻⁶. CeO_2 has been shown to be a particularly effective catalyst, in part due to the redox potential of the Ce^{4+}/Ce^{3+} couple, as well as its resistance to chemical and photocorrosion, and strong light absorption in the UV region (absorption edge \sim 385–400 nm). Unfortunately the large band gap (\sim 3.2 eV) limits further application of CeO_2^{7-8} . From the AM1.5 spectrum, it can be seen that UV light only composes \sim 3–5% of the photon flux reaching the earth's surface, while around 45% is in the visible light range. Thus it would be hugely beneficial to extend the light absorption of CeO_2 to the visible light region⁹, and as such, many methods have been investigated to shift the absorption onset by doping or coupling with other materials^{10,11}.

Doping with metal ions, specially Fe^{3+} was chosen in this research as it is believed to facilitate the Fenton reaction, which help improve the photocatalytic activity by producing the very powerful hydroxyl radicals (OH*) as oxidizers^{12,13}.

The separation of catalyst from the waste water suspension is another major problem which limits practical application of nanomaterials 14,15 . Dense films lack the high surface area required for effective catalysis. Therefore, the CeO₂ nanoparticles are fabricated to form a mesoporous film. The doctor blade deposition technique was employed for film preparation to well adhered, high surface, porous films with a good photocatalytic activity 16,17 .

In this study, kinetics studies of the photocatalytic degradation of MO over Fe-doped CeO₂ films were investigated. Improved electron-hole separation and charge transfer process mechanism was also proposed based on the obtained XPS and UV-vis DRS results. Also the formation of OH* in the photocatalytic process has been confirmed via photoluminescence spectroscopy.



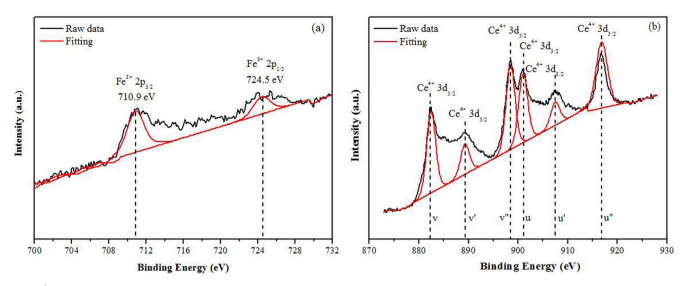


Figure 1 | The XPS spectra of 5.00 mol% Fe-doped CeO_2 (a) Fe 2p and (b) Ce 3d.

XPS analysis was carried out in order to characterise the valence states of both cerium and iron (5.00 mol% Fe-doped CeO₂). The two main peaks of Fe 2p in Figure 1a were observed at 710.9 and 724.5 eV of $2p_{3/2}$ and $2p_{1/2}$ respectively, which corresponds with the Fe³⁺ species in Fe₂O₃^{18,19}. For the Ce 3d XPS spectrum, the typical six peaks of Ce 3d are presented in Figure 1b. The peaks denoted as V (882.5 eV), V (889.3 eV), V (898.5 eV), U (901.0 eV), U (907.5 eV), and U (916.8 eV) were attributed to Ce⁴⁺ state^{20,21}.

Each of the XRD patterns in Figure 2 indexed as CeO_2 , with a cubic fluorite structure (JCPDS 3403940)²². No other peaks fromeither metallic iron or Fe_xO_y were found in the XRD analysis. Samples with higher doping revealed a shift in the peak associated with the 111 direction (located at $2\theta=28.3899$ for the undoped sample), which shifted towards higher values. The calculated crystallite size and lattice parameter decreased slightly with increasing iron content. This may be explained by larger Ce^{4+} ions (0.101 nm) being replaced by smaller Fe^{3+} ions (0.064 nm), thus decreasing the lattice parameters, unit cell volume and crystallite size²³ (Supplementary Information 1).

To obtain the band gap energies of the Fe–doped CeO $_2$ films, Schuster-Kubelka-Munk absorption function ((\propto tv) $^{1}/n$) was plotted against the photon energy (h ν) according to equation (1) 24 .

$$(\propto h\nu)^{1/n} = A(h\nu - E_g) \tag{1}$$

where A is a proportionality constant, h is Planck's constant, v is the frequency of vibration (hence hv is photon energy), and \propto is an absorption coefficient. The value of n depends on the type of optical transition of the semiconductor (n=2 for indirect transition)²⁵. The approximated band gap can then be determined from the straight line x-intercept as shown in Figure 3.

The kinetics of MO decolourisation are presented in Figure 4, and are found to follow pseudo-first order reaction as shown in equation (2)²⁶:

$$-\ln(C/C_O) = kt \tag{2}$$

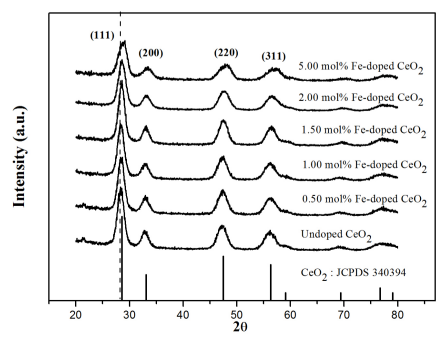


Figure 2 | XRD patterns of the undoped and Fe-doped CeO₂ films fabricated by the doctor blade technique.



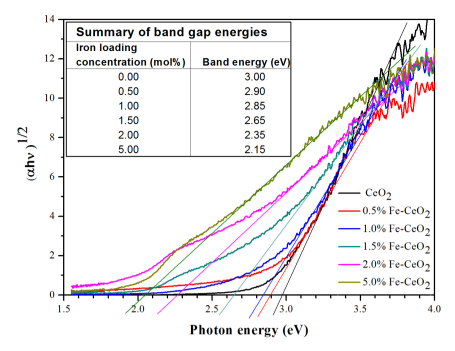


Figure 3 | Schuster-Kubelka-Munk absorption function of Fe-doped CeO₂ films with an inset showing the band gap energies of catalyst films.

where k is the apparent rate constant (min⁻¹), C_0 is the initial concentration of dye and C is the concentration of dye at time (t). As the dye concentration in these experiments remains in the regime where the Beer-Lambert law holds, the concentration can be substituted by the dye absorbance at a given wavelength (typically the peak absorption).

Iron modified CeO_2 films clearly presented superior photocatalytic activity towards the degradation of MO in pH 5 conditions compared to undoped CeO_2 , with 1.50 mol% Fe doping providing the most rapid dye degradation. BET analysis (Supplementary Information 2) showed that the 1.50 mol% Fe–doped CeO_2 film possessed the highest specific surface area, and hence more active sites are available for the MO degradation reaction²⁷.

Another important real world consideration for photocatalytic systems is long-term stability. The 1.50 mol% Fe-doped CeO_2 was employed to degrade MO solution (pH5) for 2 h before the films was removed and regenerated (heated to $250^{\circ}C$ for 120 min) and process repeated (10 cycles in total). The material displayed good stability with regards to photocatalytic performance, with less than 10% decreased from its initial activity during photodegradation process (Figure 5).

The observation of photoluminescence (PL) was employed to explain the chance of electron-hole pair recombination in catalyst. Since the PL emission spectra was reported from the recombination of excited electrons and holes^{28,29}, thus a higher PL intensity indicated a higher recombination rate of electron-hole pairs under light irra-

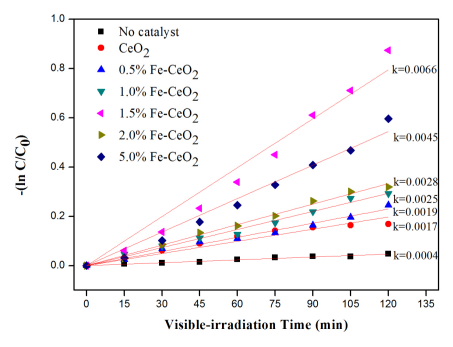


Figure 4 | The kinetics plots for pseudo first order reaction of MO decolourisation at pH 5.



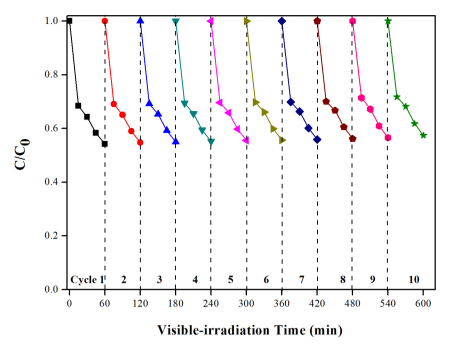


Figure 5 | Cycling runs of MO decolourisation using 1.50 mol% Fe-doped CeO₂ film at pH 5 of MO solution.

diation. In Figure 6, undoped CeO_2 presented the highest intensity, which indicated that undoped CeO_2 provided the fastest recombination rate of electrons and holes. After the iron ions were introduced to CeO_2 , the intensity of the PL emission gradually decreased, which confirmed that the doping with iron ions could delay the recombination rate of photogenerated charge carriers.

Moreover, the detection of hydroxyl radical (OH*) in photocatalytic reactions have been reported via the reaction of terephthalic acid (TA) with OH*, and formed the fluorescent 2-hydroxy terephthalate (TA–OH*) as a reaction product, which can be detected by fluorescence spectroscopy³⁰. In this study, the 1.50% Fe-doped CeO₂ catalyst was employed to detect the generated OH* on the surface. As

shown in Figure 7, the PL intensity at approximately 425 nm of TA–OH* increased in the first hour of photocatalytic degradation process and kept constant with increasing time after this one hour, which means that the catalyst presented the highest performance in the first hour of light irradiation.

Discussion

Based on the optical band gap energy values determined above (Figure 3) and equation (3) and (4), below, the conduction band (CB) and valence band (VB) potentials of CeO₂ can be calculated³¹:

$$E_{cb}(CeO_2) = \chi (CeO_2) - E^C - 1/2 Eg$$
 (3)

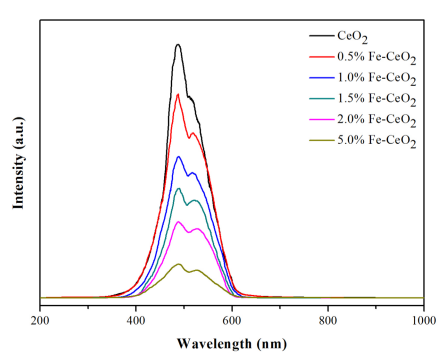
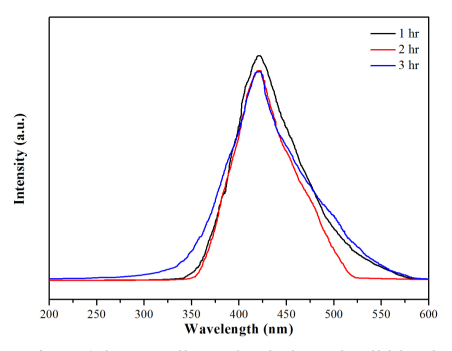


Figure 6 | PL spectra of CeO₂ at differrent amounts of iron doping level.



 $Figure \ 7 \ | \ Fluorescence \ spectra \ of \ a \ TA-OH \ \ \ 'solution \ generated \ by \ 1.50 \ mol\% \ Fe-doped \ CeO_2 \ under \ visible \ light \ irradiation.$

$$E_{vb}(CeO_2) = E_g - E_{cb}(CeO_2)$$
 (4)

Where χ is the absolute electronegativity of the semiconductor (χ is 5.56 eV for CeO₂)³²,

 E^{C} is the scaling factor relating the hydrogen electrode scale (NHE) to absolute vacuum scale (AVS) (\sim 4.5 eV ν s. AVS for 0 V ν s. NHE)³² and Eg is the band gap energy of CeO₂ (3.00 eV). The calculated CB and VB potentials of CeO₂ are -0.44 and 2.56 eV, respectively.

It is proposed the schematically in Figure 8 that the photogenerated electron-hole pairs are able to be separated for into trap states in the doped material. The presence of Fe^{3+} , as shown in XPS analysis (Figure 1a) may act as electron acceptor (from Fe^{3+} to Fe^{2+}) and/or hole donor (from Fe^{3+} to Fe^{4+}) to facilitate charge carrier localisation and hence prolonged separation by trapping at energy levels close to the conduction or valence bands, respectively³³. Therefore, the Fe^{3+} doping could be effective in producing materials that (1) delay electron–hole recombination, thereby increasing the lifetime of the elec-

tron-hole separation³⁴ as confirmed in PL results and (2) support the charge carrier transfer to the catalyst surface.

In summary, Fe-doped CeO₂ catalysts with different iron doping concentrations have successfully been fabricated to produce photocatalytic films by a doctor blade technique. Under visible light irradiation, Fe³⁺ doping can significantly increase the photocatalytic activity of CeO₂ towards the degradation of MO, with the best efficiency obtained from a 1.50 mol% iron doping. The enhanced photocatalytic performance upon doping with iron could be ascribed to both the red shift in the light adsorption range and an increased specific surface area, shown in the BET analysis. The presence of Fe³⁺ observed from XPS analysis may contributed to the Fe³⁺/Fe²⁺ and Fe⁴⁺/Fe³⁺ additional levels in the CeO₂, and a decrease in band gap energy as a consequence. The PL results confirmed that the iron doping could be effective in delay electron–hole recombination, thereby increasing the lifetime of the electron–hole separation. Since the hydroxyl radical is known to be the key species in photo-

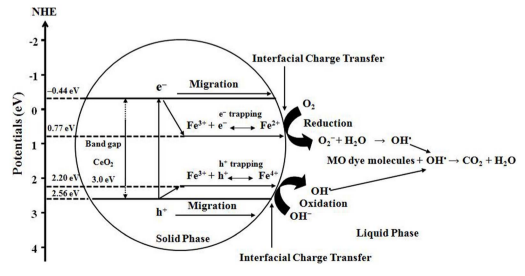


Figure 8 | Proposed mechanism for the photoexcited electron–hole separation and transport processes at the Fe-doped CeO_2 interface under visible light irradiation.



catalytic reactions, thus the PL results proved that the OH adduct TA-OH was formed during the photocatalytic reaction with the presence of catalyst.

Methods

Catalysts powder preparation. Undoped CeO_2 and Fe-doped CeO_2 powders were prepared by the combination of homogeneous precipitation and impregnation methods. Cerium (III) nitrate hexahydrate ($Ce(NO_3)_3$.6H₂O) and ferric (III) nitrate nonahydrate ($Fe(NO_3)_3$.6H₂O) were used as precursors. Firstly, a 20 mM of $Ce(NO_3)_3$.6H₂O in 100 mL of 80% ethylene glycol was prepared and stirred at 50°C until fully dissolved. During this time the colour of solution changed from purple to yellow. Addition of ammonia causes the solution to go purple once again, however after the 12 h aging a turbid yellow suspension was produced, which was then washed with deionised water, then ethanol and then dried in a vacuum oven at 70°C for 24 h. Finally, the prepared CeO_2 samples were calcined at 500°C for 1 h. Fe-doped CeO_2 with different amounts of iron concentration (0.50–5.00 mol%) were prepared by impregnation of undoped CeO_2 powders suspensionin water, with the addition of appropriate amount of $Fe(NO_3)_3$.9H₂O while continuous stirring for 2 h. The sample was then filtered, washed and dried at 70°C overnight and calcined at 500°C for 1 h.

Catalysts film preparation. Films of both doped and undoped CeO_2 were prepared by doctor blade technique as described elsewhere³⁵. The obtained film was dried in air at 50°C for about 30 min, and further annealed at 500°C for 2 h in order to remove Triton X-100 binder. The films thickness was \sim 5.0 \pm 0.2 μ m as measured by profilometer (Supplementary Information 3).

Film Characterisation. The films thickness was found to be around 5.0 \pm 0.2 μm from Profilometer measurement (Veeco Dektak150). The optical properties of the prepared samples were obtained from a photospectrometer (Shimadzu UV-3600) with integrating sphere attachment. The point of zero charge of catalyst powders was measured with the Zeta potential analysis (Malvern, Zetasizer-NanoS90). The crystalline phase of catalyst film was determined by glancing angle X-ray diffraction (GAXRD, Philips X'pert Materials). The porosity and specific surface area (SSA) were determined using Brunauer, Emmett, Teller (BET) analysis of nitrogen adsorptiondesorption data (Quantachrome Autosorp 1 MP). The chemical composition and oxidation state of all materials were studied by X-ray photoelectron spectroscopy (XPS) using Mg X-ray source (MgKα, Kratos Axis Ultra DLD). The binding energy of the adventitious carbon (C 1s) line at 285 eV was used for calibration and the position of other peaks was corrected according to the position of the C 1s signal. The thickness cross-section of the films was measured by scanning electron microscopy (FESEM, JSM-6335F, JEOL). The fluorescence spectra of the generated TA-OH were measured on the fluorescence spectrophotometer at 425 nm (Avantes, AvaSpec-2048 TEC-USB2-2); fluorescence was induced by excitation at 345 nm.

Photocatalytic testing for the degradation of dyes. Photocatalytic degradation processes under visible irradiation were investigated using undopedCeO2 and FedopedCeO2 films through the decolourisation of MO solution with an initial dye concentration of 0.025 mM at pH 5. The adsorption efficiency of MO in dark at different pH value are explained in Supplementary Information 4 and 5. The photoreactor system employed with a halogen lamp (50 watt power) providing a light intensity of $\sim\!185$ mW cm $^{-2}$. Decolourisation of dye solutions at different time intervals was determined by measuring the change in absorbance maxima of MO at 464 nm via UV-vis spectrophotometer (Shimadzu UV-1800).

Determination of reactive species (hydroxyl radical, OH'). In order to measure the relative concentration of hydroxyl radicals, the terephthalic acid (TA) fluorescence method was employed in this study owing to the TA can react with OH' forming highly fluorescent 2-hydroxyterephthalic acid (TA–OH'). The mixture of TA solution was prepared from 5×10^{-4} molar of TA and 6×10^{-3} molar of NaOH in DI water. Then, 0.25 g of the 1.50% Fe-doped CeO $_2$ catalyst was dispersed in 50 mL of the TA aqueous solution. The solution was collected at every one hour during the irradiation procedure in order to estimate the generated TA–OH', which can be detected by fluorescence spectroscopy at 425 nm³ 6,37 .

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Author contributions

D.C. analysed the data and wrote the manuscript. S.P. and B.I. supervised the project, had given valuable advices on the proceeding of this work, and revised the manuscript. A.N. and J.C. designed the concept and the experiment method of the research. S.U. supported the characterisation of the samples. N.W. had provided valuable suggestions on the selection of supplementary information. All authors discussed the results and commented on the manuscript at all stages.

Additional information

Supplementary information accompanies this paper at http://www.nature.com/ scientificreports

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