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Enhanced water splitting through two-step photoexcitation by sunlight using tantalum/nitrogen-codoped rutile titania as a water oxidation photocatalyst†

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Rutile TiO₂ codoped with tantalum and nitrogen (TiO₂:Ta,N) was assessed as a water oxidation photocatalyst for Z-scheme water splitting driven by visible light. This material was prepared by thermal ammonolysis of TiO₂:Ta with dry NH₃ at 773 K, while samples of the oxide precursor were synthesized using a microwave-assisted solvothermal technique, applying various conditions. The photocatalytic activity of the TiO₂:Ta,N during water oxidation to O₂ from an aqueous FeCl₃ solution was found to be greatly affected by the synthesis parameters. The rate of O₂ evolution was increased upon increasing the level of Ta doping in conjunction with a highly-crystallized TiO₂:Ta precursor resulting from applying a higher temperature during synthesis of the oxide. IrO₂ loading of the TiO₂:Ta,N photocatalyst also improved the O₂ evolution activity. The optimized IrO₂/TiO₂:Ta,N photocatalyst was applied to a Z-scheme water splitting system in combination with Ru/SrTiO₃:Rh and in the presence of redox mediators (Fe^{3+/2+} or [Co(bpy)₃]^{3+/2+}). Under AM1.5G simulated sunlight, this system exhibited a maximum solar-to-hydrogen energy conversion efficiency of 0.039%, which was nearly twice as high as the previously reported system with the use of RuO₂/TiO₂:Ta,N (0.021%).

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Introduction

The photocatalytic splitting of water into H₂ and O₂ is one potential means of producing H₂ using solar energy.^{1–6} Semiconductors capable of absorbing visible light have been extensively studied over the last half century as photocatalysts for water splitting, because visible light accounts for approximately one half of the energy contained in sunlight.^{1–6} Various reaction schemes have been proposed for water splitting. Among these, the two-step photoexcitation system referred to as the Z-scheme,

which uses two different semiconductor photocatalysts, is the most promising because it requires lower energy photons and allows the separation of the H₂/O₂ mixture that is generated during the water splitting reaction.^{4–6} In the Z-scheme process, the two semiconductors catalyze H₂ and O₂ evolution, while a redox shuttle reagent mediates electron transfer between the two materials (Scheme 1). Many semiconductors, including doped metal oxides, solid solutions and mixed anion compounds, have been assessed as visible light water-splitting photocatalysts applicable to either water reduction or oxidation in Z-scheme water splitting.^{4–18}

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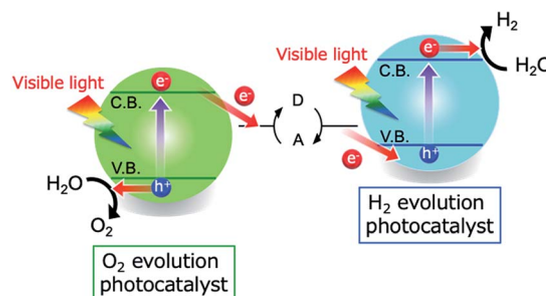
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Scheme 1 A diagram showing a Z-scheme water splitting system consisting of two different semiconductors and a redox shuttle mediator.



Doping foreign elements into a metal oxide is a conventional approach to obtaining new properties that might assist in water splitting, such as visible light absorption^{12,19,20} and high carrier concentrations.^{21,22} As an example, doping of a metal oxide with nitrogen can narrow the band gap of the oxide as a result of substitution of oxide ions with nitride ions, because the valence band potential of the oxide is shifted negatively through the contribution of the N 2p orbitals.^{19,20} The formation of carriers due to foreign element doping^{21,22} or defect formation²³ in a metal oxide also improves the photocatalytic activity of the oxide, especially with regard to the H₂ or O₂ evolution half reactions. For these reasons, carefully designed metal oxide photocatalysts based on doping have been shown to function as efficient H₂ or O₂ evolution photocatalyst in Z-scheme water splitting.^{4–13,15–18}

Our group has recently reported a highly active visible-light-responsive photocatalyst, tantalum/nitrogen codoped rutile titanium dioxide (TiO₂:Ta,N), that promotes the oxidation of water to O₂.^{7,24} This photocatalyst can be synthesized *via* the thermal ammonolysis of rutile TiO₂:Ta which, in turn, is produced using a microwave-assisted solvothermal approach. A solar-to-hydrogen energy conversion efficiency (STH) of 0.021% has been obtained using RuO₂-modified TiO₂:Ta,N and Ru-loaded SrTiO₃:Rh as water oxidation and reduction photocatalysts, respectively, in the presence of an Fe³⁺/Fe²⁺ redox shuttle mediator. Transient absorption spectroscopy has demonstrated that the codoped tantalum ions in TiO₂:Ta,N suppress the defect formation that is otherwise associated with nitrogen doping, thereby prolonging the lifetime of photo-generated free electrons. Therefore, pre-doping of the rutile TiO₂ with Ta is essential to the efficient functioning of this catalyst. That is, the precursor oxide (rutile TiO₂:Ta) plays an important role in synthesizing the final TiO₂:Ta,N material.

Microwave-assisted hydro/solvothermal synthesis is an effective approach to the preparation of nanostructured oxide materials, including photocatalyst powders.^{25–28} Compared to conventional heating, in which the container is heated from the outside, microwave radiation produces a rapid increase in the internal temperature of the substrate because the microwaves are directly absorbed by solvent, with the temperature rise varying based on the dielectric dissipation factor of the solvent. This rapid heating can lead to interesting phenomena, such as increased reaction rates and selective phase formation.^{29,30} While microwave-assisted synthesis has been widely applied to the production of various metal oxides,^{25,27–30} no systematic investigation of the preparation of doped metal oxides such as TiO₂:Ta has been reported.

The photocatalytic activity of a semiconductor can be promoted by surface modification with metal/metal oxide nanoparticles, acting as so-called cocatalysts.^{2,31,32} These cocatalysts facilitate the separation of excited charge carriers^{2,32} and can also serve as redox reaction sites.^{2,32} The modification of semiconductor photocatalysts with cocatalysts is especially effective (and even indispensable in some cases) with regard to Z-scheme water splitting in aqueous solutions containing redox shuttle mediators. This is because the cocatalyst improves the selectivity for forward reactions while suppressing undesirable

backward reactions.^{5,6,32–34} As an example, Domen *et al.* reported that a RuO₂ cocatalyst on TaON promotes both the reduction of IO₃[−] and the oxidation of water.³³ Abe *et al.* developed a RuO₂ cocatalyst that is active during the selective reduction of IO₃[−] and also improves O₂ evolution when used as a co-modifier for a cobalt-based water oxidation cocatalyst.³⁴ Because the effects of loading cocatalysts on rutile TiO₂:Ta,N photocatalysts have not yet been investigated in detail, the activity increases provided by such cocatalysts can likely still be significantly improved.

Based on this background, the present work attempted to improve the photocatalytic water oxidation activity of rutile TiO₂:Ta,N by refining the synthesis conditions for the semiconductor itself and by optimizing the cocatalyst loading. Rutile TiO₂:Ta was synthesized by a microwave-assisted solvothermal approach under various conditions and the resulting materials were used as precursors to obtain TiO₂:Ta,N by thermal nitridation with dry NH₃ gas. The effects of the synthesis conditions applied during the solvothermal reaction on the photocatalytic activity of the TiO₂:Ta,N during water oxidation were investigated in detail. Optimizing the preparation conditions as well as the cocatalyst was found to improve the performance of this catalyst during solar-driven Z-scheme water splitting.

Experimental

Preparation of photocatalysts

Ta-doped rutile TiO₂ powders were synthesized *via* a microwave-assisted technique previously reported in the literature.⁷ In this method, a quantity of Ti(OiPr)₄ (Kanto Chemical, >97.0%, 19.8–19.86 mmol) and an appropriate volume of Ta(OEt)₅ (Sigma-Aldrich, 99.98%, 0.14–0.2 mmol) were added to *n*-hexane (Kanto Chemical, >96.0%, 50 mL) containing HCl (Kanto Chemical, 35–37%, 7 mL) after which the reaction solution was placed in an autoclave and heated at a specific temperature (373–423 K) for 2 h using microwave radiation (400 W). The resulting precipitate was washed with ethanol (Kanto Chemical, >99.5%) and water several times, and then dried at 343 K in an oven. Nitridation of the TiO₂:Ta powders was performed by annealing at 773 K under an NH₃ flow (Sumitomo Seika Chemicals, >99.9995%, 10 mL min^{−1}) for 30 min in an alumina tube connected to a mass-flow controller.

Cocatalyst-loaded samples were prepared by a previously reported impregnation method.³³ In each case, the photocatalyst powder was initially dispersed in an aqueous solution of the desired metal precursor with stirring. The precursor was Co(NO₃)₂·6H₂O (FUJIFILM Wako Pure Chemical Corporation, >98.0%), RuCl₃·*n*H₂O (Furuya Metal, Ru 41.35 wt%), Na₃RhCl₆·*n*H₂O (Mitsuwa Chemicals, Rh 15.3 wt%) or Na₂IrCl₆·6H₂O (Kanto Chemical, >97.0%). This was followed by evaporation of the solvent and heating at 573 K for 1 h in air.

The SrTiO₃:Rh photocatalyst was obtained by a previously reported hydrothermal method.^{15,35} In a typical synthesis, Sr(OH)₂·8H₂O (Kanto Chemical, >96.0%, 22 mmol), TiO₂ (JRC-TIO-10, The Catalysis Society of Japan, 19.6 mmol) and Rh(NO₃)₃ (Kanto Chemical, >80%, 0.5 mmol) were mixed in 50 mL of water, after which the mixture was transferred to



a Teflon-lined stainless-steel autoclave and heated at 433 K for 42 h. The resulting precipitate was washed with hot water and room-temperature water, then dried at 363 K in an oven. The hydrothermal product was subsequently mixed with an additional amount of $\text{Sr}(\text{OH})_2 \cdot 8\text{H}_2\text{O}$ (1.4 mmol) and heated at 1273 K for 10 h in air. The resulting material was washed with water and dried at 363 K. Finally, the Ru cocatalyst was deposited using a photodeposition method.^{13,18} In this process, the $\text{SrTiO}_3\text{:Rh}$ (0.3 g) was stirred in an aqueous methanol (Kanto Chemical) solution (100 mL, 10 vol%) containing RuCl_3 (3 wt% vs. $\text{SrTiO}_3\text{:Rh}$) for 3 h and then washed with water several times. The powder was collected and then dispersed in a 10 vol% aqueous methanol solution, followed by irradiation at $\lambda > 400$ nm for 3 h to obtain Ru-loaded $\text{SrTiO}_3\text{:Rh}$ (Ru/ $\text{SrTiO}_3\text{:Rh}$). The Ru cocatalyst loading on the $\text{SrTiO}_3\text{:Rh}$ was approximately 2.2 wt%, as determined by inductively coupled plasma-mass spectrometry (ICP-MS) analysis.

Characterization of photocatalysts

The prepared samples were studied by powder X-ray diffraction (XRD; Rigaku, MiniFlex 600; Cu $K\alpha$), UV-visible diffuse reflectance spectroscopy (DRS; JASCO, V-670), transmission electron microscopy (TEM; JEOL, JEM-2010F) and X-ray photoelectron spectroscopy (XPS; Shimadzu, ESCA-3400). The binding energies were corrected with respect to the position of the C 1s peak (285.0 eV) for each sample. The chemical compositions were determined using ICP optical emission spectroscopy (ICP-OES; Agilent Technologies, 5100 VDV ICP-OES) and ICP-MS (Agilent Technologies, 7700x). Nitrogen concentrations were assessed using an elemental micro analyzer (J-Science, JM10). The Brunauer-Emmett-Teller (BET) surface areas of the specimens were measured using a BELSORP-mini (MicrotracBEL) at liquid nitrogen temperature (77 K).

STEM observations

Scanning transmission electron microscopy (STEM) images were acquired using an electron microscope (Thermo Fisher Scientific, Titan cubed) equipped with spherical aberration correctors (CEOS) and an energy dispersive X-ray spectroscopy (EDS) apparatus, at an acceleration voltage of 300 kV. The convergence semiangle of the incident probe was 18 mrad, with an incident probe current of 18 pA. The detector collection semiangle was from 45.8 to 200 mrad for high-angle annular dark-field (HAADF).

Transient absorption spectroscopy

Transient absorption spectroscopy data were acquired using laboratory-built spectrometers described previously.^{36,37} Briefly, $\text{TiO}_2\text{:Ta,N}$ samples were excited by 450 nm laser pulses from a Nd:YAG laser (Continuum, Surelite I, 6 ns duration, 1.0 mJ, 1 Hz) and the transmittance and reflectance for each specimen were determined below and above 6000 cm^{-1} , respectively, using mid-IR ($6000\text{--}1000\text{ cm}^{-1}$; MoSi coil), near-IR ($10\,000\text{--}6000\text{ cm}^{-1}$; halogen lamp, 50 W) and visible ($25\,000\text{--}10\,000\text{ cm}^{-1}$; halogen lamp, 50 W) irradiation. The probe light was dispersed by the spectrometer while the monochromatized mid-IR, near-IR and visible light was detected by an MCT

detector (Kolmar), an InGaAs detector and a Si photodiode, respectively. The time resolution of this spectrometer was limited to the range of 1–2 μs as a result of the incorporation of an AC-coupled amplifier (Stanford Research Systems, SR560) in the mid-IR and near-IR regions, and to approximately 4 μs , due to stray light associated with the pump pulse and/or short-lived strong emission from the sample in the visible region.

Photocatalytic reactions

Photocatalytic O_2 evolution reactions were conducted at room temperature using a top-irradiation type cell that was connected to a closed gas circulation system made of glass. In each trial, a 50 mg sample was dispersed in 140 mL of a 10 mM aqueous AgNO_3 solution or a 1 mM aqueous FeCl_3 (Kanto Chemical, >99.0%) solution (100 mL). In trials in which the O_2 evolution reaction was conducted in the co-presence of Fe^{2+} ions, FeCl_2 (FUJIFILM Wako Pure Chemical Corporation, 99.0–102.0%) was also added to the reactant solution. In each trial, the reaction solution was first outgassed using a vacuum pump, after which Ar gas was introduced into the reaction system to a pressure of approximately 5 kPa, and the solution was irradiated with a 300 W Xe lamp (Cermax, PE300BF) operating at an output current of 20 A. The light from the lamp was collected using a CM-1 cold mirror and then passed through an L-42 cut-off filter ($\lambda > 400$ nm) prior to reaching the sample. The gases evolved during the reaction were analyzed by gas chromatography (GL Sciences, GC-3200 with a thermal conductivity detector and Ar as the carrier gas).

Z-scheme overall water splitting trials were conducted in a similar manner. Quantities of the cocatalyst-modified $\text{TiO}_2\text{:Ta,N}$ (50 mg) and Ru/ $\text{SrTiO}_3\text{:Rh}$ (25 mg) powders were dispersed in an aqueous FeCl_3 (100 mL, pH 2.4) or tris(2,2'-bipyridyl) cobalt(II) sulfate ($[\text{Co}(\text{bpy})_3]^{2+}$, 0.5 mM) solution (100 mL, pH 3.8). The cobalt complex was synthesized according to a previously reported method prior to these trials.¹⁶ The pH of the reaction solution was adjusted to the desired value with H_2SO_4 , after which the solution was outgassed and irradiated ($\lambda > 400$ nm). In the case of experiments employing simulated sunlight, a HAL-320 solar simulator (Asahi Spectra) served as the light source.

The solar-to-hydrogen energy conversion efficiency (STH) was calculated according to the equation:

$$\text{STH}(\%) = \frac{R_{\text{H}} \Delta G^0}{PS} \times 100,$$

where R_{H} , ΔG^0 , P and S are the rate of H_2 evolution (mol s^{-1}) during Z-scheme water splitting, the standard Gibbs free energy of formation for water ($237 \times 10^3\text{ J mol}^{-1}$), the intensity of the simulated sunlight (100 mW cm^{-2}) and the irradiation area (16 cm^2), respectively.

Results and discussion

Effects of Ta doping level

Previous work demonstrated that doping of rutile TiO_2 with Ta followed by nitridation at the appropriate temperature was



necessary to obtain highly active $\text{TiO}_2\text{:Ta,N}$,⁷ but the optimal doping amount was not determined. For this reason, we initially performed a detailed investigation of the effects of the Ta doping amount employed during the microwave-assisted solvothermal synthesis on the physicochemical and photocatalytic properties of the $\text{TiO}_2\text{:Ta,N}$. In the first such trials, the microwave heating temperature was fixed at 423 K.

Fig. 1 shows XRD patterns for samples with different Ta doping levels, before and after nitridation. The full width at half maximum (FWHM) values for the diffraction peaks at 36° are summarized in Table S1.† Prior to nitridation, each sample produced a single-phase diffraction pattern corresponding to that expected for rutile TiO_2 , with no evidence of impurities (Fig. 1a). The diffraction peaks were observed to become sharper (*i.e.*, the FWHM decreased) after annealing under a flow of NH_3 gas (Fig. 1b), indicating that the nitridation promoted crystal growth. In addition, the XRD pattern for the sample containing 1.5 mol% Ta exhibited peaks associated with anatase TiO_2 after ammonolysis. The positions of the diffraction peaks generated by the oxide precursors changed very little after doping, even at higher Ta doping levels. This lack of change can likely be attributed to the relatively low amount of Ta that was added and/or to the minimal difference in the ionic radii of Ta^{5+} and Ti^{4+} in a six coordination environment.³⁸ In addition, no noticeable change in peak positions could be identified after nitridation.

Fig. 2 presents the UV-visible DRS data for the same set of samples. Each of the oxide materials exhibited an absorption edge at approximately 405 nm, which remained almost unchanged regardless of the Ta doping amount (Fig. 2a). In contrast, the optical properties of the semiconductors were found to change following nitridation (Fig. 2b). Specifically, an absorption band attributed to electron transitions from N 2p to Ti 3d orbitals^{19,20} appeared in the region of 400–500 nm following nitridation, although the absorption band at approximately 400 nm (assigned to the band gap of the host $\text{TiO}_2\text{:Ta}$) also remained. The degree of visible light absorption also varied with respect to the amount of Ta added, although there was no clear trend in these variations.

Photocatalytic O_2 evolution reactions were conducted using the as-prepared $\text{TiO}_2\text{:Ta,N}$ powders, modified with 0.8 wt% of

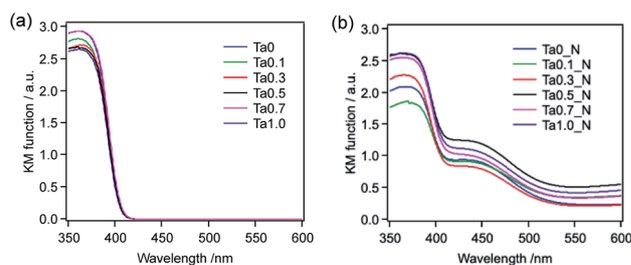


Fig. 2 UV-visible DRS spectra of $\text{TiO}_2\text{:Ta}$ prepared with different levels of Ta doping (a) before and (b) after nitridation at 773 K.

a RuO_2 cocatalyst, in aqueous 1 mM FeCl_3 solutions under visible light irradiation ($\lambda > 400$ nm). As noted in the Introduction, the RuO_2 acted as a bifunctional cocatalyst for reduction/oxidation reactions.³¹ The photocatalytic activity during O_2 evolution was found to sharply increase with increasing Ta doping concentration up to 0.5 mol%, while the undoped sample did not generate O_2 under the present conditions (Fig. 3). The O_2 evolution rate reached a maximum at 0.5 mol% Ta and essentially plateaued with further increase in

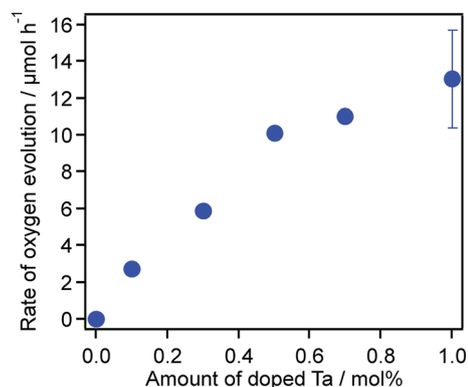


Fig. 3 Photocatalytic O_2 evolution activity for $\text{TiO}_2\text{:Ta,N}$ samples as a function of Ta doping level. Reaction conditions: catalyst = 50 mg (cocatalyst = RuO_2 at 0.8 wt%); reactant solution = aqueous FeCl_3 (1 mM, 100 mL); light source = Xe lamp (300 W) with a cold mirror (CM-1) and a cutoff filter (L42).

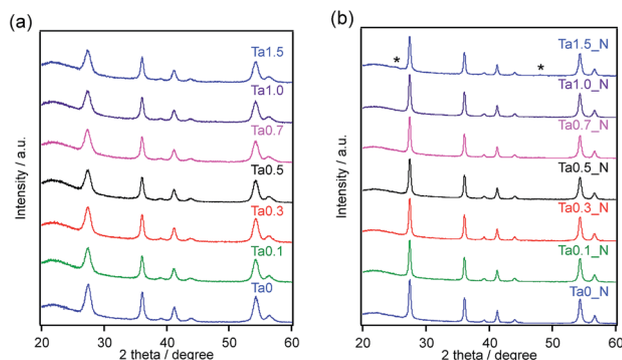


Fig. 1 XRD patterns for $\text{TiO}_2\text{:Ta}$ prepared with different levels of Ta doping: (a) before and (b) after nitridation at 773 K. Asterisks indicate diffraction peaks attributed to anatase TiO_2 .

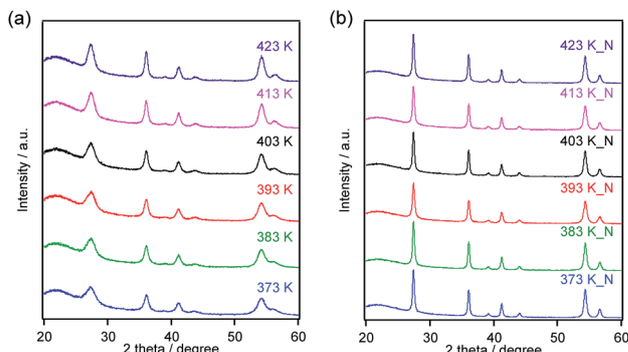


Fig. 4 XRD patterns for $\text{TiO}_2\text{:Ta}$ specimens prepared at different temperatures (a) before and (b) after nitridation at 773 K.



Table 1 Physicochemical properties of TiO₂:Ta,N specimens prepared at different temperatures

Entry	Heating temperature/K	FWHM ^a /degree		Specific surface area/m ² g ⁻¹		Nitrogen content in TiO ₂ :Ta,N/wt%
		Before nitridation	After nitridation	Before nitridation	After nitridation	
1	373	0.801	0.296	80	12	0.11
2	383	0.702	0.305	73	22	0.10
3	393	0.723	0.351	41	15	0.12
4	403	0.634	0.294	45	12	0.11
5	413	0.616	0.318	86	14	0.13
6	423	0.550	0.314	37	10	0.17

^a Full width at half maximum for the (101) peak ($2\theta = 36.0$ degree) in the XRD patterns.

the doping level. On the basis of these results, we concluded that a Ta doping amount of 0.5 mol% or more is required to permit TiO₂:Ta,N to exhibit high photocatalytic activity during visible-light water oxidation. In the following sections, during analysis of the microwave synthesis, the Ta doping was therefore fixed at 0.7 or 1.0 mol%.

Effects of microwave heating temperature

Because the microwave treatment was expected to greatly affect the TiO₂:Ta product, we investigated the effects of applying different temperatures while keeping all other synthesis parameters constant and employing a Ta concentration of 1.0 mol%. Fig. 4a shows XRD patterns for TiO₂:Ta specimens prepared using different temperatures, while Table 1 summarizes the FWHM values for the peaks at 36°. All the products produced single-phase diffraction patterns attributable to rutile TiO₂. The diffraction peaks became more intense and the FWHM decreased with increasing heating temperature, indicating an improvement in crystallinity.

Fig. 5 shows the particle morphology, as observed by TEM. While the morphology of the specimens heated at 383 and 423 K was similar (Fig. 5a and b), a portion of the latter sample evidently underwent crystal growth, forming larger particles with sizes of several hundred nanometers (Fig. 5c). This growth likely resulted from rapid, localized heating in response to the microwave radiation, and could also be related to irregular changes in the specific surface areas of the products (Table 1).

The optical properties of the oxide precursors were examined by means of UV-visible DRS. As shown in Fig. S1,† each specimen showed an absorption edge at approximately 405 nm, although those samples having higher surface areas had

absorption edges that were blue-shifted, most likely due to quantum size effects.

The subsequent nitridation process varied the physicochemical properties of the materials dramatically. Fig. 4b shows XRD patterns for the nitrated products, using the same set of oxide materials. Each specimen produced a single-phase diffraction pattern assignable to a rutile structure after nitridation. The FWHM values for the peaks were reduced compared to those for the oxides prior to nitridation (Table 1). These results indicate that crystal growth occurred upon heating at 773 K under a flow of NH₃, in agreement with the TEM observations (Fig. S2†). This crystal growth during nitridation was accompanied by a decrease in the specific surface area, to approximately 10–20 m² g⁻¹. Interestingly, the evident variations in the crystallinity, specific surface area and morphology between the oxide precursors disappeared completely after nitridation.

Nitridation also changed the optical properties of these materials, and Fig. 6 presents the UV-visible DRS spectra of the same series of nitrated samples. As noted above, an absorption band in the range of 400–500 nm appeared after nitridation. While the extent of visible light absorption varied irregularly with respect to the microwave heating temperature, the sample processed at 373 K showed significantly less absorption in the visible region of the spectrum. The nitrogen content was measured by elemental analysis of the same set of nitrated samples, and found to be in the range of 0.1–0.2 wt% (Table 1).

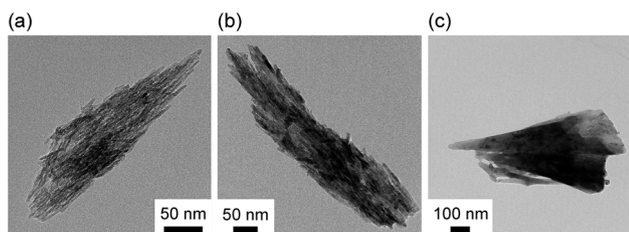


Fig. 5 TEM images of TiO₂:Ta (1.0 mol%) samples prepared at (a) 373 and (b and c) 423 K.

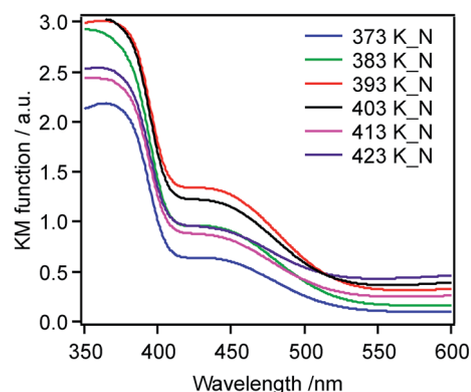


Fig. 6 UV-visible DRS spectra for TiO₂:Ta,N specimens made from oxide precursors (TiO₂:Ta) prepared at different temperatures.



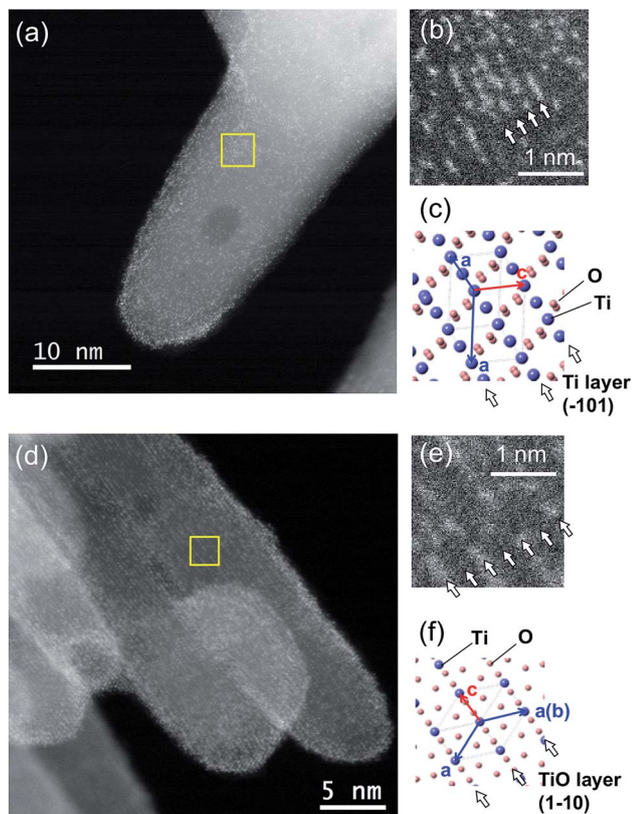


Fig. 7 HAADF-STEM images of $\text{TiO}_2\text{:Ta,N}$ powder (Ta 0.7 mol%). (a and d) HAADF-STEM images and (b and e) enlarged views of the regions indicated by the yellow squares in (a and d), respectively. (c and f) Crystal structure models of TiO_2 from the [121] and [111] directions.

A detailed investigation of the physicochemical state of the Ta dopants was performed so as to ascertain the occupation sites, the actual amounts and the oxidation states. Fig. 7 shows HAADF-STEM images of a $\text{TiO}_2\text{:Ta,N}$ powder with a nominal Ta concentration of 0.7 mol%. The brighter spots in these images correspond to Ta ions because the signal intensity in HAADF

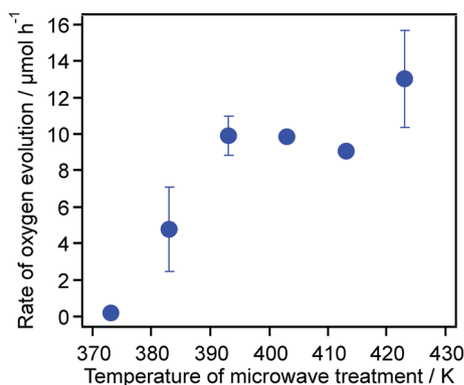


Fig. 8 Photocatalytic O_2 evolution activity for $\text{TiO}_2\text{:Ta,N}$ samples as a function of the temperature used to prepare the precursor. Reaction conditions: catalyst = 50 mg (cocatalyst = RuO_2 at 0.8 wt%); reactant solution = aqueous FeCl_3 (1 mM, 100 mL); light source = Xe lamp (300 W) with a cold mirror (CM-1) and a cutoff filter (L42).

imaging is approximately proportional to Z^2 (where Z represents the atomic number).³⁹ Here, the Ta sites are seen to be located in $\text{Ti}\{101\}$ and $\text{TiO}\{110\}$ layers. These sites are uniformly distributed throughout the $\text{TiO}_2\text{:Ta,N}$ particles. The concentration of Ta was determined using EDS and ICP-OES. The Ta atom concentrations determined by these techniques were 0.6 and 0.9 mol% (*vs.* Ti), respectively, both of which are close to the nominal value (0.7 mol%).

The surface oxidation state of the Ta atoms was assessed using XPS, and Fig. S3† presents the Ta 4f XPS spectrum of $\text{TiO}_2\text{:Ta,N}$ (Ta: 0.7 mol%) powder. Two peaks are evident, at 28.1 and 26.3 eV, which can be assigned to Ta $4f_{5/2}$ and $4f_{7/2}$ orbitals in the Ta_2O_5 .⁴⁰ It is known that the position of the Ta 4f photoelectron signal is sensitive to the local environment around Ta atoms,⁴¹ and these results show that the Ta atoms in the $\text{TiO}_2\text{:Ta,N}$ were in the Ta(v) oxidation state even after thermal ammonolysis.

Fig. 8 plots the O_2 evolution rates for the $\text{TiO}_2\text{:Ta,N}$ samples as a function of the temperature applied during the microwave synthesis of the oxide precursors. The O_2 evolution rate was dramatically accelerated with increasing temperature up to 393 K, beyond which it remained almost unchanged.

Factors affecting the photocatalytic activity of $\text{TiO}_2\text{:Ta,N}$

The results for the photocatalytic reactions indicate that the activity of $\text{TiO}_2\text{:Ta,N}$ during O_2 evolution was significantly affected by the preparation parameters (*i.e.*, the Ta doping amount and the microwave treatment temperature), as shown in Fig. 3 and 8. There were no noticeable differences in the structural and physicochemical properties among $\text{TiO}_2\text{:Ta,N}$ materials with different Ta doping levels (Fig. 1 and Table S1†). In addition, the changes observed in the UV-visible DRS spectra were minimal and somewhat irregular (Fig. 2). However, in the case of photocatalytic O_2 evolution reactions in aqueous FeCl_3 solutions, the activity was dramatically changed with increasing amount of Ta. In prior work, transient absorption data revealed that Ta doping is necessary to obtain a highly active photocatalyst, because the Ta suppresses the formation of oxygen defects in the $\text{TiO}_2\text{:Ta,N}$, thus prolonging the lifetimes of photogenerated free and/or shallowly trapped electrons.⁷ Considering the results of the present study, a Ta doping level of at least 0.5 mol% is likely sufficient for the suppression of defects.

The temperature applied during the microwave treatment was also found to be a very important factor affecting the photocatalytic activity (Fig. 8). Changing the temperature resulted in different degrees of crystallinity in the $\text{TiO}_2\text{:Ta}$ precursors (Fig. 4a and Table 1), such that the photocatalytic activity of the $\text{TiO}_2\text{:Ta,N}$ during O_2 evolution was enhanced with increasing temperature. While the activity of these materials did not exhibit an obvious correlation with visible light absorption (Fig. 6) or surface area (Table 1), the O_2 evolution rate was found to increase as the FWHM of the XRD peaks for the oxide precursors decreased (Table 1). That is, the use of more highly crystalline $\text{TiO}_2\text{:Ta}$ tends to yield higher photocatalytic activity in the resulting $\text{TiO}_2\text{:Ta,N}$.



Transient absorption spectroscopic analyses of samples synthesized at different microwave treatment temperatures were performed to determine the lifetime of photogenerated free and/or shallowly trapped electrons. Two TiO₂:Ta,N samples that showed different activities (383 and 423 K samples) were selected. As shown in Fig. S4,[†] the lifetimes for these two specimens were almost the same, indicating that the excited carrier dynamics in both materials were similar. These equivalent excited carrier lifetimes as well as the similarities in the other physicochemical properties (*e.g.*, crystallinity, specific surface area and light absorption) suggest that the primary factors affecting the O₂ evolution activity are related to surface rather than bulk characteristics. Interestingly, when Ag⁺ was used as an electron acceptor, the sample heated at 383 K showed a higher rate of O₂ evolution (35.1 μmol h⁻¹) than the 423 K sample (21.0 μmol h⁻¹), as shown in Fig. S5,[†] which was opposite to the results obtained using Fe³⁺ as the electron acceptor (Fig. 8; 383 K, 4.8 μmol h⁻¹ and 423 K, 13.2 μmol h⁻¹).⁴² The reduction of Ag⁺ to Ag is irreversible, while the Fe³⁺ to Fe²⁺ reduction is reversible, meaning that there is no competing reductant in a reactant solution containing aqueous AgNO₃. In addition, Fe²⁺ can be photo-oxidized to Fe³⁺, thereby reducing the rate of the forward water oxidation reaction. We therefore propose that the higher activity for the 423 K sample during O₂ evolution from aqueous FeCl₃ is attributable to suppression of the reverse reaction (*i.e.*, the oxidation of Fe²⁺ to Fe³⁺) and/or enhancement of the reduction of Fe³⁺.

On the basis of these results, we concluded that the appropriate choice of the oxide precursor for the TiO₂:Ta,N is very important for obtaining a highly active TiO₂:Ta,N photocatalyst. The critical factors are the Ta doping level (which should be greater than 0.5 mol%) and high crystallinity, which can be obtained by heating at temperatures above 393 K.

Effect of cocatalyst loading on the photocatalytic activity of TiO₂:Ta,N

Using the optimized TiO₂:Ta,N, we subsequently investigated the effect of cocatalyst loading on the photocatalytic water oxidation activity. Table 2 provides the O₂ evolution rates for aqueous FeCl₃ solutions containing TiO₂:Ta,N (Ta: 0.7 mol%) modified with various metal oxide cocatalysts (0.8 wt% each), loaded by an impregnation method. Each of these metal oxide

Table 2 The effects of modifying TiO₂:Ta,N with various metal oxide cocatalysts on the rate of O₂ evolution in aqueous FeCl₃ solutions under visible light ($\lambda > 400$ nm)^a

Entry	Cocatalyst	Rate of O ₂ evolution/μmol h ⁻¹
1	None	0.8
2	CoO _x	2.6
3	RuO ₂	11.0
4	RhO _x	2.5
5	IrO ₂	14.1

^a Reaction conditions: catalyst = 50 mg; reactant solution = aqueous FeCl₃ (1 mM, 100 mL); light source = Xe lamp (300 W) with a cold mirror (CM-1) and a cutoff filter (L42).

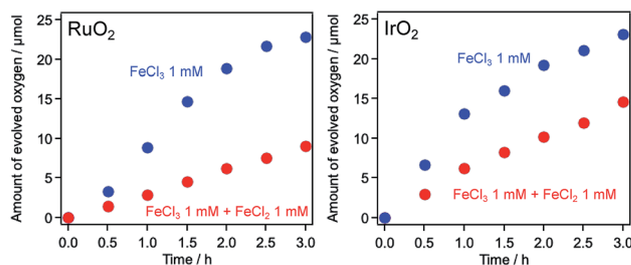


Fig. 9 Time courses for oxygen evolution over RuO₂-loaded and IrO₂-loaded TiO₂:Ta,N under visible light irradiation ($\lambda > 400$ nm) in aqueous solutions containing Fe³⁺ and Fe²⁺ ions. Reaction conditions: catalyst = 50 mg (cocatalyst as noted = 0.8 wt% each); reactant solution = aqueous FeCl₃ (1 mM, 100 mL) with or without FeCl₂ (1 mM); light source = Xe lamp (300 W) with a cold mirror (CM-1) and a cutoff filter (L42).

cocatalysts facilitated the O₂ evolution process as compared to the unloaded sample. The IrO₂-modified sample showed the highest O₂ evolution rate, with stable performance during a 15 h irradiation trial (Fig. S6[†]). O₂ evolution over the IrO₂/TiO₂:Ta,N was found to cease when 25 μmol of O₂ had been generated, which corresponds to the amount expected from the stoichiometry of a reaction involving the four-electron oxidation of water and the one-electron reduction of Fe³⁺ (100 μmol).

During the O₂ evolution reaction from an aqueous FeCl₃ solution, photo-oxidation of Fe²⁺ to Fe³⁺ can occur, competing with the forward water oxidation reaction. As discussed in the Introduction, it is important to suppress such reverse reactions, because the photo-oxidation of Fe²⁺ is thermodynamically more favorable than water oxidation.⁴³ The behavior of the IrO₂/TiO₂:Ta,N photocatalyst during the O₂ evolution reaction suggests that the performance of this material is relatively unaffected by the reverse reaction involving Fe²⁺ to Fe³⁺ oxidation. To further investigate the effect of IrO₂ loading on photocatalytic O₂ evolution from aqueous FeCl₃ solutions, O₂ evolution reactions were performed in the presence of Fe²⁺ ions. As shown in Fig. 9, the O₂ evolution activity for a RuO₂-loaded sample was comparable to that for an IrO₂-loaded sample in an aqueous FeCl₃ solution. In contrast, in an aqueous solution containing both Fe³⁺ and Fe²⁺ ions, the activity for the former material was lower than that for the latter. Iridium oxide is known to be an excellent water oxidation catalyst (or cocatalyst) with regard to the generation of O₂,⁴⁴ and so loading of IrO₂ on

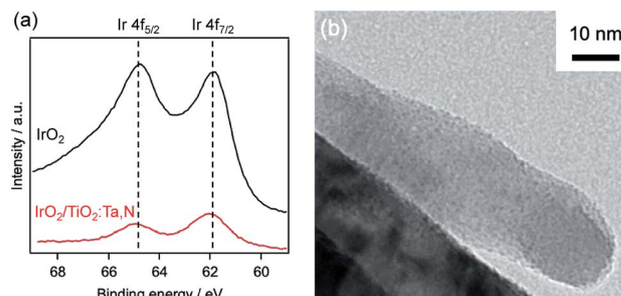


Fig. 10 Characterization of 1.0 wt% IrO₂-loaded TiO₂:Ta,N. (a) Ir 4f XPS spectrum and (b) TEM image.



the $\text{TiO}_2\text{:Ta,N}$ likely resulted in efficient water oxidation sites even in the presence of Fe^{2+} . This would explain the higher activity for the IrO_2 -loaded sample as compared to the material containing RuO_2 .

The physicochemical characteristics of the most active IrO_2 -loaded material were further studied, and the electronic states of the IrO_2 loaded on the $\text{TiO}_2\text{:Ta,N}$ were investigated by XPS. Fig. 10a shows the Ir 4f XPS spectrum of IrO_2 -loaded $\text{TiO}_2\text{:Ta,N}$ powder with a nominal Ta level of 0.7 mol%, along with reference data for pure IrO_2 . The spectrum exhibits two major peaks at binding energies of 64.9 and 62.0 eV, which are consistent with the Ir 4f_{5/2} and 4f_{7/2} peaks in the IrO_2 reference spectrum.⁴⁵ This result clearly indicates that the Ir on the $\text{TiO}_2\text{:Ta,N}$ was present as IrO_2 .

As shown in Fig. 10b, TEM observations indicated that the IrO_2 deposits were uniformly dispersed on the $\text{TiO}_2\text{:Ta,N}$ surface, in the form of nanoparticles with sizes of approximately 1 nm. These particles were readily distinguishable due to the contrast in the images resulting from the different electron densities for Ir and Ti (or Ta).

Z-scheme water splitting using $\text{TiO}_2\text{:Ta,N}$ as an O_2 evolution photocatalyst

IrO_2 -loaded $\text{TiO}_2\text{:Ta,N}$ was subsequently applied as the water oxidation component in a Z-scheme water splitting system in combination with $\text{SrTiO}_3\text{:Rh}$ as a H_2 evolution photocatalyst and in the presence of an $\text{Fe}^{3+/2+}$ redox couple. $\text{SrTiO}_3\text{:Rh}$, which was originally developed by Kudo *et al.*, is a promising photocatalyst for H_2 evolution under visible light.^{15,16,46} The XRD pattern and UV-visible DRS spectrum of the $\text{SrTiO}_3\text{:Rh}$ synthesized in this work are shown in Fig. S7.†

Table 3 presents the rates of H_2 and O_2 evolution from an aqueous FeCl_3 solution containing IrO_2 -loaded $\text{TiO}_2\text{:Ta,N}$ (Ta: 0.7 wt%) and Ru-loaded $\text{SrTiO}_3\text{:Rh}$. These rates were strongly accelerated in the presence of a small amount of FeCl_3 (0.2 mM, entry 2). At higher FeCl_3 concentrations, the activity decreased

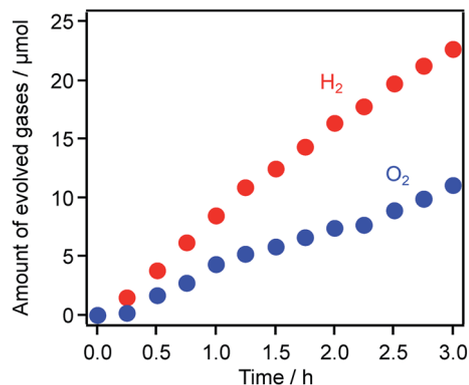


Fig. 11 Time dependence of solar-driven H_2 and O_2 evolution from mixtures of $\text{IrO}_2\text{/TiO}_2\text{:Ta,N}$ (50 mg) and $\text{Ru/SrTiO}_3\text{:Rh}$ (25 mg) dispersed in an aqueous FeCl_3 solution (100 mL, 0.2 mM) under simulated sunlight irradiation (AM1.5G, 100 mW cm^{-2}).

gradually (entries 3–5), possibly due to the reverse reaction of Fe^{3+} photoreduction. Unmodified $\text{TiO}_2\text{:Ta,N}$ also acted as a water oxidation photocatalyst with $\text{Ru/SrTiO}_3\text{:Rh}$ in the presence of the $\text{Fe}^{3+/2+}$ redox mediator, producing nearly stoichiometric amounts of H_2 and O_2 (entry 6). The water splitting activity in an aqueous FeCl_3 solution was, however, improved upon loading with the IrO_2 cocatalyst, by a factor of approximately 4 (entries 2, 7–9). These data again confirm the promotional effect of the IrO_2 cocatalyst. In addition, the activity was increased upon increasing the concentration of the IrO_2 cocatalyst to 1.0 wt%, beyond which it began to drop. Stoichiometric water splitting into H_2 and O_2 under visible light irradiation was also achieved using a combination of $\text{IrO}_2\text{/TiO}_2\text{:Ta,N}$ and $\text{Ru/SrTiO}_3\text{:Rh}$ in aqueous solutions containing $[\text{Co}(\text{bpy})_3]^{2+}$ (Fig. S8†), which has been reported to be another effective redox mediator for Z-scheme water splitting.¹⁶

Finally, Z-scheme water splitting was conducted under simulated sunlight (AM1.5G, 100 mW cm^{-2}). As shown in Fig. 11, because H_2 and O_2 were generated in a nearly stoichiometric ratio, it is clear that this system exhibited solar energy conversion through overall water splitting to store chemical energy in the form of H_2 . The STH was calculated to be 0.039%, which is 1.9 times higher than the previously reported value for a system using RuO_2 -loaded $\text{TiO}_2\text{:Ta,N}$ (0.021%).⁷ In addition, the STH value recorded in this work is comparable to (or a bit higher than) that achieved using a combination of $\text{Ir/CoO}_x\text{/Ta}_3\text{N}_5$ and $\text{Ru/SrTiO}_3\text{:La/Rh}$ (0.037%), the highest recorded STH for a Z-scheme system based on an (oxy)nitride photocatalyst as the O_2 evolution component.⁴⁷

Conclusions

A microwave-assisted solvothermal approach was applied to the synthesis of Ta-doped rutile TiO_2 , which was then subjected to thermal nitridation with dry NH_3 gas to yield Ta/N-codoped rutile TiO_2 ($\text{TiO}_2\text{:Ta,N}$). The use of a highly crystalline $\text{TiO}_2\text{:Ta}$ precursor, which was obtained by microwave treatment at higher temperatures, was important for obtaining $\text{TiO}_2\text{:Ta,N}$

Table 3 Effects of iridium oxide modification and redox shuttle mediator on water splitting reaction under visible light ($\lambda > 400 \text{ nm}$)^a

Entry	Concentration of $\text{FeCl}_3\text{/mM}$	Amount of loaded $\text{IrO}_2\text{/wt\%}$	Rate of gas evolution/ $\mu\text{mol h}^{-1}$		
			H_2	O_2	$\text{H}_2\text{/O}_2$ ratio
1	0	0.8	4.3	2.1	2.0
2	0.2	0.8	23.7	12.7	1.9
3	0.5	0.8	21.9	12.0	1.8
4	1.0	0.8	18.8	10.2	1.8
5	2.0	0.8	16.2	8.8	1.8
6	0.2	0	8.0	3.8	2.1
7	0.2	1.0	29.9	15.5	1.9
8	0.2	1.2	21.5	10.9	2.0
9	0.2	1.5	17.4	9.1	1.9

^a Reaction conditions: catalyst = $\text{IrO}_2\text{/TiO}_2\text{:Ta,N}$, 50 mg and $\text{Ru/SrTiO}_3\text{:Rh}$, 25 mg; reactant solution = aqueous FeCl_3 (100 mL, pH 2.4); light source = Xe lamp (300 W) with a cold mirror (CM-1) and a cutoff filter (L42).



that exhibited high photocatalytic activity during visible-light water oxidation. It was also found that a minimum level of Ta dopant (here, greater than 0.5 mol%) was necessary to activate the full potential of the TiO₂:Ta,N photocatalyst, most likely because of the associated improvement in the lifetime of photogenerated mobile electrons. While TiO₂:Ta,N showed activity during water oxidation in the presence of Fe³⁺ as a reversible electron acceptor, modification with an IrO₂ nanoparticle cocatalyst resulted in a marked activity enhancement, by a factor of 17. The IrO₂-loaded TiO₂:Ta,N photocatalyst could be applied to a Z-scheme water splitting system combined with Ru/SrTiO₃:Rh in aqueous FeCl₃ (or [Co(bpy)₃]²⁺) solutions under visible light. The maximum solar-to-hydrogen energy conversion efficiency reached 0.039%, which was comparable to values obtained from state-of-the-art Z-scheme systems incorporating (oxy)nitride-based materials as water oxidation photocatalysts.

Although the visible light absorption capability of this doped oxide photocatalyst was inferior to that of (oxy)nitrides, we believe that carefully designed doped oxides have significant potential as photocatalysts for visible light water splitting, as clearly demonstrated in this work. Because there are a large number of metal oxides that could be used as host materials for the fabrication of doped photocatalysts, there is still much capacity for the development of more active doped photocatalysts for solar-driven water splitting. This possibility is currently under investigation in our laboratory.

Author contributions

K. M. designed the project. S. N. conducted most of the experiments and analysis, together with K. M. and M. I., J. J. M. V. and A. Y. performed the transient absorption spectroscopic analyses, D. L. conducted the TEM observations, and K. Y. and K. K. conducted the STEM observations. S. N. and K. M. wrote a draft of the manuscript. All authors discussed and provided comments on the experiments and the manuscript during preparation.

Conflicts of interest

There are no conflicts to declare.

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