RSC Advances



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Cite this: RSC Adv., 2015, 5, 100123

Hydrothermal synthesis of rhodium-doped barium titanate nanocrystals for enhanced photocatalytic hydrogen evolution under visible light†

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Rhodium-doped barium titanate (BaTiO₃:Rh) nanocrystals were prepared by hydrothermal (HT) method followed by post-heating, and examined as photocatalysts for H₂ evolution from an aqueous methanol solution under visible light (λ > 420 nm). The photocatalytic activity was dependent on synthetic parameters such as precursor ratio and using precursor materials, post-heating conditions, which have influence on crystallinity, compositional homogeneity and the generation of Ba defects in BaTiO3:Rh, as well as the oxidation state of Rh in the material. Experimental results highlighted that suppressing the generation of Ba defects during the preparation process and improving the crystallinity both contributed to higher photocatalytic activity. The most active material prepared by the HT method outperformed similar analogues prepared by the polymerized complex method and a conventional solid state reaction method.

Received 28th September 2015 Accepted 13th November 2015

DOI: 10.1039/c5ra20044i

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Introduction

Hydrogen has drawn significant attention as a clean energy carrier, which outputs high energy without releasing harmful byproducts. The development of a heterogeneous photocatalyst, which can produce H₂ via water splitting with visible light, is an important topic in chemistry, and various materials including doped metal oxides, solid solutions, (oxy)nitrides and (oxy) sulfides have been developed as photocatalysts. 1-3

Metal oxide photocatalysts are in general stable under irradiation, and some have achieved very high quantum yields of several tens of percent under UV irradiation. However, there are only a few metal oxides that can reduce and oxidize water under visible light.2 Therefore, development of a visible-lightresponsive oxide photocatalyst that can produce H2 from water with visible light is of importance to construct a nonsacrificial Z-scheme water splitting system.^{2,4}

Rh doping is one of the most effective methods that enables one to produce a visible-light-responsive photocatalyst, as reported by Kudo and co-workers.5 Our group has recently reported that Rh-doped BaTiO₃ having a cubic perovskite structure prepared by solid-state reaction (SSR) method and polymerized complex (PC) method, showed photocatalytic activities for H2 evolution from water under visible light irradiation.6 The BaTiO3:Rh photocatalyst produces H2 without satisfactory, and further improvement in activity is required. Hydrothermal (HT) method is an environmental benign,

noticeable degradation in Z-scheme overall water splitting for as

long as 30 hours. Unfortunately, however, the activity is not

effective method for preparing perovskite materials that show high photocatalytic activity.7,8 Kominami et al. have developed a new HT synthetic method (HyCOM; hydrothermal crystallization in organic media) to obtain TiO2 nanocrystals, which exhibited very high photocatalytic activity for many kinds of reactions including H₂/O₂ evolution and the mineralization of acetic acid.9,10 Domen et al. have applied a HT method for the synthesis of NaTaO3 perovskite.7 According to that study, NaTaO₃ prepared by the HT method showed higher photocatalytic activity for overall water splitting than a SSR-derived analogue. For the synthesis of a doped metal oxide, Kudo et al. recently reported that photocatalytic activity of Rh-doped SrTiO₃ prepared by HT method for H₂ evolution is comparable to that obtained by the PC method.8 While HT method has been well studied for the synthesis of various pristine metal oxides not only for photocatalytic applications but also for others, 11 however, systematic investigation on the preparation parameters of a doped metal oxide has not been made so far, and the structure-activity relationship has not been clarified enough.

Base on these backgrounds, this study aimed at preparing Rh-doped BaTiO₃ by a HT method under various synthesis conditions, in order to improve the photocatalytic activity for H₂ evolution. The influence of synthesis parameters and postheating treatment on the photocatalytic activity is discussed on the basis of structural characterization.

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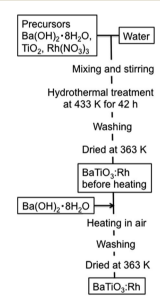
† Electronic supplementary information (ESI) available: Additional characterisation and reaction data. See DOI: 10.1039/c5ra20044i

2. Results and discussion

2.1. Precursor effects in the HT process on the formation of $BaTiO_3$:Rh

Scheme 1 shows the preparation procedure of $BaTiO_3$:Rh by a HT method and subsequent post-heating. The HT synthesis was conducted at 433 K for 42 h in a Teflon-lined stainless autoclave containing an aqueous suspension of TiO_2 dissolved with Ba^{2+} and Rh^{3+} ions without any other additives such as surfactants. The concentration of Rh was 2.0 mol% unless otherwise stated.

We first investigated effects of ${\rm TiO_2}$ precursor on the formation of ${\rm BaTiO_3:Rh}$ using four different ${\rm TiO_2}$ particles, which are P25 (a mixture of anatase, rutile and amorphous ${\rm TiO_2}$ nanoparticles), 12 nanosized anatase (JRC-TIO-8), nanosized rutile (JRC-TIO-6), and submicron-size anatase (purchased from Kanto Chemicals Co.). Structural properties of these ${\rm TiO_2}$ precursors are summarized in Table 1, and typical TEM images are shown in Fig. S1.†



Scheme 1 Preparation scheme of BaTiO₃:Rh by a hydrothermal method, followed by post-heating.

Table 1 Physicochemical properties of TiO₂ used as the precursor in this work

${ m TiO}_2$	Particle size/nm	Specific surface area/m² g ⁻¹	Crystal phase
P25	10-50	51	Mixture of anatase, rutile and amorphous
Submicron anatase (Kanto	50-120	16	Anatase
Chemicals Co.) Nano anatase (JRC-TIO-8)	\sim 5	231	Anatase
Nano rutile (JRC-TIO-6)	30-50	81	Rutile

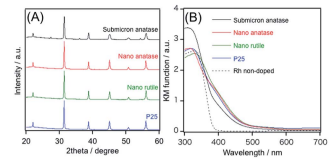


Fig. 1 (A) XRD patterns and (B) UV-visible diffuse reflectance spectra of products obtained from different TiO_2 precursors without post-heating. An excess amount of $\text{Ba}(\text{OH})_2 \cdot \text{8H}_2\text{O}$ (10 mol%) was added in the HT process.

Fig. 1A shows X-ray diffraction (XRD) patterns of products obtained from different ${\rm TiO_2}$ precursors without post-heating. A 10 mol% excess barium hydroxide was added into the mixture of starting materials in all cases. All of the prepared samples, except for the sample using submicron-size anatase, showed single-phase diffraction patterns attributed to cubic perovskite BaTiO₃. No significant difference with respect to FWHM and peak positions could be identified. However, in the sample prepared from submicron-size anatase, a small amount of unreacted ${\rm TiO_2}$ phase was observable.

Fig. 1B shows UV-visible diffuse reflectance spectra of the same set of the samples, along with a spectrum of undoped BaTiO₃ for comparison. As reported previously, undoped BaTiO₃ shows no absorption band in visible light region.⁶ However, the Rh-doped materials just after HT synthesis were yellow powder, exhibiting a visible light absorption band at around 500 nm. This absorption can be assigned to Rh³⁺ species in BaTiO₃:Rh.⁶ In the sample prepared from submicron-size anatase, the absorption band at 600–700 nm, assignable to Rh⁴⁺ species, is more pronounced, and the 500 nm absorption band is slightly weaker compared to others. This indicates that the densities of Rh⁴⁺ and Rh³⁺ species in the sample prepared from submicron-size anatase are greater and smaller than those in others.

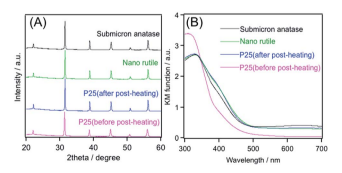
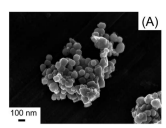


Fig. 2 (A) XRD patterns and (B) UV-visible diffuse reflectance spectra of hydrothermally synthesized BaTiO $_3$:Rh using P25, nano-size rutile and submicron-size anatase, further subject to post-heating at 1173 K. An excess amount of Ba(OH) $_2$ ·8H $_2$ O (10 mol%) was added in the HT process.

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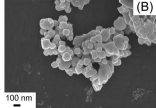


Fig. 3 SEM images of a hydrothermally synthesized $BaTiO_3$:Rh using P25 as the precursor. (A) Before and (B) after post-heating at 1173 K.

After post-heating of a hydrothermally synthesized BaTiO3:-Rh in air at 1173 K for 7 h, XRD patterns became stronger and narrower (Fig. 2A). This result indicates that crystallization of the product proceeds during post-heating. The crystal growth was also evident from the result of SEM observations (Fig. 3), accompanied with a decrease in the specific surface area from 8 to 4 m² g⁻¹. In addition, no noticeable difference was observed in all of the post-heated samples, as shown in Fig. 2A. However, the remaining anatase phase observed in the submicron-size anatase sample before post-heating (see Fig. 1A) disappeared completely. Fig. 2B shows diffuse reflectance spectra of the same set of the samples. Upon post-heating, absorption at longer wavelengths due to Rh4+ species was pronounced only when submicron-anatase was employed as the precursor. On the other hand, the spectral feature of samples prepared using P25 and nanosized rutile resembles each other. The details of post-heating will be described in the latter section.

2.2. Effects of excess addition of barium hydroxide on the formation of $BaTiO_3$:Rh

Because addition of an excess amount of Ba2+ ions in the HT process was expected to affect the production of BaTiO₃:Rh, we investigated this by altering the initial amount of barium hydroxide. XRD and DRS analyses showed that no noticeable difference in the HT products could be identified, even when the HT synthesis was conducted using a stoichiometric mixture (i.e., the ratio of Ba/(Ti + Rh) = 1.00), compared to the material prepared with an excess barium hydroxide (Fig. S2†). An exception is that the Rh4+ absorption band in the stoichiometric sample after post-heating (green line in Fig. S2†) is slightly more pronounced, compared to the excess Ba material (black line). As listed in Table 2, however, ICP-MS analysis indicated a clear difference in the chemical composition. It showed that the Ba/ (Ti + Rh) ratio in the stoichiometric sample was 0.93 (entry 1), smaller than the ideal value of 1.00, while the excess Ba sample had the Ba/(Ti + Rh) ratio of 1.00 (entry 2), almost equal to unity. This means that the stoichiometric sample contained Ba defects in the crystal lattice more or less.

Post-heating of a hydrothermally synthesized metal oxide at elevated temperatures will in general result in further crystallization and/or crystal growth, which may contribute to an enhanced photocatalytic activity. The hydrothermally synthesized samples were then subject to post-heating in the presence of an excess amount of barium hydroxide (5.0 mol% excess).

Table 2 Results of ICP-MS analyses^a

Entry	HT synthesis with an excess Ba(OH) ₂	Post-heating with an excess Ba(OH) ₂	,
1	No	_	0.93
2	Yes	_	1.00
3	No	Yes	1.00
4	Yes	Yes	1.06
5	No	No	0.91
6	Yes	No	0.99

^a The reproducibility of the ratio measured by ICP-MS was within 0.01–0.02. In each case, HT synthesis was made using P25 as the precursor in the presence of an excess $Ba(OH)_2 \cdot 8H_2O$ (10 mol%).

Again XRD measurements showed that there was no noticeable difference between the two after post-heating (Fig. S3 \dagger). Also, no noticeable change among these samples could be identified in SEM as well (Fig. S4 \dagger). However, ICP-MS analysis indicated that the Ba/(Ti + Rh) ratio of the sample, which was prepared using stoichiometric precursors during HT process, was 1.00 (entry 3). It means that the Ba defects were compensated to a certain extent by addition of barium hydroxide in the post-heating step. For the sample prepared using excess barium hydroxide during the HT process, the Ba/(Ti + Rh) ratio of the sample was increased from 1.00 to 1.06, as expected. XPS measurement showed that the surface atomic ratio of Ba/(Ti + Rh) was approximately 1.5, much higher than the Ba/(Ti + Rh) ratio determined by ICP-MS, meaning that Ba species are segregated in the surface region.

Without addition of an excess barium hydroxide in the post-heating step, the Ba/(Ti + Rh) ratio of BaTiO₃:Rh tended to decrease upon post-heating at 1173 K, regardless of whether or not an excess amount of barium hydroxide was added in the HT process (entries 5 and 6). These results suggest that post-heating at elevated temperatures might generate Ba defects in BaTiO₃:Rh, and the addition of barium hydroxide in the post-heating step can suppress the defect formation. More detailed investigation on the post-heating effect will be shown in the latter section.

2.3. Photocatalytic activities of BaTiO₃:Rh prepared using different precursors

Then we evaluated photocatalytic activities of the synthesized materials using different precursors. For these samples, postheating was made in static air at 1173 K for 7 h in the presence of an additional barium hydroxide (5.0 mol% excess). As mentioned later, post-heating was indispensable for obtaining high activity. As listed in Table 3, photocatalytic activity of BaTiO₃:Rh depended strongly on the precursors employed in the HT process. In all cases, Pt (0.5 wt%) was loaded onto BaTiO₃:Rh by an *in situ* photodeposition method in order to promote hydrogen evolution catalysis, and it was found that 0.5 wt% Pt loading was the best to perform the reaction. TEM observation indicated that Pt nanoparticles of 3–5 nm in size were distributed on the BaTiO₃:Rh surface with some

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Table 3 Photocatalytic activities of BaTiO₃:Rh prepared using different TiO_2 precursors for H₂ evolution from an aqueous methanol solution under visible light ($\lambda > 420 \text{ nm}$)^a

Entry	${ m TiO}_2$	Specific surface area/ m ² g ⁻¹	Ratio of Ba/ (Ti + Rh) in the starting mixture of the HT process	H ₂ evolved (3 h)/μmol
1	P25	4	1.10	42.3
2	Nano anatase	9	1.10	38.7
3	Nano rutile	6	1.10	18.7
4	Submicron	2	1.10	13.3
	anatase			
5	P25	4	1.00	22.0

^a Reaction conditions: catalyst, 100 mg (0.5 wt% Pt photodeposited *in situ*); 10 vol% aqueous methanol solution 100 mL; light source, xenon lamp (300 W) fitted with a cold mirror (CM-1) and a cutoff filter (L42); reaction vessel, Pyrex top-irradiation type. Post-heating was made for all of the samples at 1173 K for 7 h in the presence of an excess amount of Ba(OH)₂·8H₂O (5.0 mol%).

aggregation (Fig. S5†). Samples prepared using P25 and nanosized anatase, which consist of nanoparticulate anatase as the main phase, showed relatively high activity (entries 1 and 2). Using nanoparticulate rutile as the precursor, however, the activity was approximately half of those recorded using P25 and nanosized anatase (entry 3), even though these XRD patterns and DRS were very similar (Fig. 2). When submicron-size anatase was used as the precursor, the activity was about 30% of that recorded using P25 and nanosized anatase (entry 4). It was thus found that TiO_2 that contains nanosized anatase as the main phase was better as the precursor for HT synthesis of BaTiO₃:Rh nanocrystals.

We also found that adding an excess amount of barium hydroxide in the HT process was important for activity-enhancement. The activity of BaTiO₃:Rh prepared using the stoichiometric amount of precursors (*i.e.*, Ba/(Ti+Rh) = 1.00) in the HT process was relatively low (entry 5), even though an additional amount of barium hydroxide was supplied in the subsequent post-heating process.

2.4. Effects of post-heating on physicochemical properties of BaTiO₃:Rh

More detailed investigation on the HT synthesis of BaTiO₃:Rh was made, focusing on the effects of post-heating temperature of the hydrothermally-synthesized BaTiO₃:Rh on physicochemical and photocatalytic properties. For these investigations, an excess barium hydroxide (10 mol%) was added in the HT process, based on the previous finding mentioned above.

Fig. 4A shows XRD patterns of BaTiO₃:Rh post-heated at different temperatures in the presence of an additional barium hydroxide (5.0 mol%). Compared to the unheated material, BaTiO₃:Rh post-heated at 873 K exhibited sharper and more intense diffraction peaks, indicating the occurrence of crystal growth upon post-heating. However, increasing the heating temperature up to 1273 K did not alter the diffraction patterns. SEM observation showed that particles of the 1173 K sample are

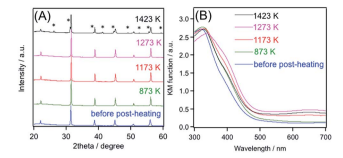


Fig. 4 (A) XRD patterns and (B) UV-visible diffuse reflectance spectra of BaTiO $_3$:Rh post-heated at different temperatures in the presence of an additional Ba(OH) $_2 \cdot$ 8H $_2$ O (5.0 mol%). The star * indicates hexagonal BaTiO $_3$.

sintered to some extent to form larger secondary particles, compared to the sample just after HT treatment, as already shown in Fig. 3. This observation is consistent with the result of XRD measurement (Fig. 4A). At 1423 K, hexagonal BaTiO₃ phase was generated, accompanied with a decrease in the diffraction peak intensity of cubic BaTiO₃ phase.

On the other hand, DRS underwent a successive change with increasing temperature; at elevated temperatures except for the sample at 1423 K, absorption at longer wavelengths due to Rh⁴⁺ species was more pronounced (Fig. 4B). This would be reasonable, because heat-treatment at elevated temperatures promoted oxidation of Rh3+ into Rh4+ in BaTiO3:Rh. We also tried to quantify the Rh⁴⁺/Rh³⁺ existence ratio in these samples by means of XPS. However, the position of the Rh3d_{5/2} peak remained before and after post-heating (Fig. S6†), different from the result of DRS measurement (Fig. 4B). It is likely that the change in the valence state of the Rh species between the two samples was too small to be detected by XPS measurements. A similar result has been obtained in our previous study on the same material prepared by the PC method.6 In that case, the Rh3d_{5/2} XPS peak of the as-prepared BaTiO₃:Rh did not undergo a change even after hydrogen reduction treatment.

2.5. Photocatalytic activities of post-heated BaTiO3:Rh

Table 4 lists photocatalytic activities of hydrothermally synthesized BaTiO₃:Rh further post-heated at different temperatures. Here an additional amount of barium hydroxide (5.0 mol% excess) was mixed with a hydrothermally synthesized BaTiO₃:Rh before post-heating. The unheated sample exhibited relatively low activity (entry 1). However, the photocatalytic activity was increased with a rise of the post-heating temperature to reach a maximum at 1273 K (entry 4), beyond which it began to drop significantly. Interestingly, even upon post-heating at suitable temperature, we could not obtain high activity without addition of an excess barium hydroxide (entry 6).

As the result of applying HT synthesis and optimizing the synthetic parameters, we could obtain an apparent quantum yield for $\rm H_2$ evolution of ca. 0.8% at 420 nm, which was 1.5 times higher than that recorded by the PC method.⁶ It should be also noted that the activity of the HT sample was about an order of magnitude higher than that of the SSR sample.⁶

Table 4 Photocatalytic activities of BaTiO $_3$:Rh post-heated at different temperatures for H $_2$ evolution from an aqueous methanol solution under visible light ($\lambda > 420 \text{ nm}$) a

Entry	Heating temperature/ K	Addition of 5.0 mol% $Ba(OH)_2 \cdot 8H_2O$ in the postheating step	Specific surface area/ m ² g ⁻¹	H ₂ evolved (3 h)/μmol
1	_	Yes	8	10.4
2	873	Yes	8	12.8
3	1173	Yes	4	42.1
4	1273	Yes	4	54.1
5	1423	Yes	3	1.9
6	1173	No	4	19.4

^a Reaction conditions: catalyst, 100 mg (0.5 wt% Pt photodeposited *in situ*); 10 vol% aqueous methanol solution 100 mL; light source, xenon lamp (300 W) fitted with a cold mirror (CM-1) and a cutoff filter (L42); reaction vessel, Pyrex top-irradiation type. HT synthesis was made using P25 as the precursor in the presence of an excess $Ba(OH)_2 \cdot 8H_2O$ (10 mol%).

2.6. Factors affecting photocatalytic activity of BaTiO₃:Rh

The results of photocatalytic reactions indicated that the activity of BaTiO3:Rh for H2 evolution was strongly dependent on the preparation parameters both in the HT process and in postheating step (Tables 3 and 4). In the HT process, an appropriate choice of TiO2 precursor was found to be very important to produce an active BaTiO3:Rh photocatalyst. As shown in Fig. 2, single-phase diffraction patterns assigned to a perovskite structure were observed for samples prepared using nanosized TiO₂, regardless of the crystal structure (anatase or rutile). On the other hand, small peaks assigned to anatase TiO2 remained in the sample prepared using submicron anatase. Although post-heating of these samples at 1173 K resulted in the production of a single-phase BaTiO3:Rh, it is clear that the sample prepared using larger anatase has compositional inhomogeneity, which could account for the relatively low activity of the material. It is most likely that the reactivity of larger TiO₂ with other precursors of Ba and Rh in the HT process was too low to produce a single-phase perovskite material.

To support this idea, we synthesized BaTiO₃:Rh in the same manner using P25 TiO₂, but with different Rh precursors of Rh₂O₃ and RhCl₃, which are respectively insoluble and soluble in water. As shown in Fig. S7,† XRD analysis showed that a single phase diffraction pattern assignable to BaTiO₃ was obtained when a water-soluble Rh precursor (*i.e.*, Rh(NO₃)₃ and RhCl₃) was used, while the Rh₂O₃ derived sample contained impurity phase. As expected, BaTiO₃:Rh prepared from Rh₂O₃ produced a trace amount of H₂, while Rh(NO₃)₃ and RhCl₃-derived ones showed appreciable activity (Fig. S8†).

It can be thus concluded that reactivity of precursors in the HT process is a critical factor that enhances the photocatalytic activity of BaTiO₃:Rh; in the present case, using nanosized TiO₂ is essential. Interestingly, however, the activity of BaTiO₃:Rh prepared using nanosized rutile TiO₂ was almost half of that derived from nanosized TiO₂ consisting of anatase as the main phase, even though there is almost no difference in XRD and

DRS (Fig. 2). During the formation of BaTiO₃:Rh from a TiO₂ aqueous suspension containing Ba²⁺ and Rh³⁺ in the HT process, it is most likely that rearrangement of constituent ions occurs because the crystal structure of TiO₂ is different from that of BaTiO₃:Rh. ICP-MS analysis showed that the rutile derived material had the Ba/(Ti + Rh) ratio of 0.96, even though the synthesis was conducted in the presence of an excess amount of barium hydroxide both in the HT process and postheating step. This result indicates that this sample contains Ba defects in its structure, which is most likely to contribute to the lower photocatalytic activity. Note here that the activity was almost independent of the specific surface area.

In addition to suitable precursors for the HT synthesis, addition of an excess amount of barium hydroxide before the synthesis was essential to obtaining $BaTiO_3$:Rh that shows higher activity. Even with addition of barium hydroxide (5.0 mol% Ba) after the HT process (that is, just before post-heating), the activity was approximately half of those prepared with excess Ba (see Table 3). ICP-MS analyses showed that the ratio of Ba/(Ti+Rh) in the HT product prepared from stoichiometric amounts of precursors was 0.93. This shortage of Ba indicates the formation of Ba defects.

In the post-heating step, both heating temperature and addition of an excess amount of barium hydroxide were important factors. The enhanced activity of BaTiO3:Rh with increasing temperature (see Table 4 and Fig. S7†) is primarily associated with the improvement in the crystallization of the material. In the temperature range up to 1273 K, where the largest increase in activity was obtained, the XRD peak became stronger and narrower (Fig. 4A) and the particles of BaTiO3:Rh grew to some extent (Fig. 3), consistent with the change in the specific surface area. Crystallization decreases the density of defects and/or structural imperfections that can act as recombination centers between photogenerated electrons and holes, resulting in an enhanced activity.2,4 Another possible cause of the enhanced activity with temperature is the generation of suitable density of Rh4+ species, as observed in UV-visible DRS (Fig. 4B). Kudo et al. claimed that during photocatalytic H₂ evolution on SrTiO₃:Rh, part of Rh⁴⁺ ions in the crystal lattice undergo photoreduction by electrons generated in the conduction band of the material.5 As the result, unstable Rh3+ ions, which are more effective for the H2 evolution than the stabilized Rh3+ ions by oxygen defects, are formed. However, further increase in calcination temperature resulted in a significant drop in activity (Table 4 and Fig. S7†). Although this might be due to the generation of Rh⁴⁺ species that do not undergo photoreduction, XRD data clearly indicated that the structure of cubic perovskite BaTiO3:Rh was destroyed upon heating at 1423 K, generating hexagonal phase (Fig. 4A). This might be another reason for the very low activity of the 1423 K sample.

The addition of an excess amount of barium hydroxide with BaTiO₃:Rh in the post-heating step was found to improve the photocatalytic activity. Even though HT synthesis was conducted in the presence of excess barium hydroxide to prevent the formation of Ba defects, it was found that addition of barium hydroxide in the post-heating process was important to

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enhancing photocatalytic activity. As mentioned earlier, ICP-MS analysis showed that some of the incorporated Ba species in BaTiO₃:Rh were lost during the post-heating process (Table 2). Therefore, the addition of barium hydroxide in the post-heating process helped to suppress defect formation, which would be essential to enhancing photocatalytic activity of BaTiO3:Rh for H₂ evolution. It indicates that post-heating with barium hydroxide resulted in the generation of Ba-rich BaTiO3:Rh, which might be more suitable for application in photocatalytic H₂ evolution.

Conclusions

BaTiO3:Rh is active for photocatalytic H2 evolution from an aqueous methanol solution under visible light irradiation. In this work, we synthesized BaTiO3:Rh by a hydrothermal method followed by post-heating, and investigated the synthetic parameters. In order to enhance the photocatalytic activity, it was important to improve the compositional homogeneity and suppress the generation of Ba defects in BaTiO3:Rh, while maintaining high crystallinity and the oxidation state of Rh(III) in the material.

Experimental 4.

4.1. Preparation of photocatalyst powders

TiO₂ powders were obtained from Japan Aerosil, Catalysis Society of Japan, and Kanto Chemicals. For the Rh source, Rh(NO₃)₃ (Kanto Chemicals, >80.0%), Rh₂O₃ (Wako Pure Chemical Industries, >98.0%) and RhCl₃·3H₂O (Wako Pure Chemical Industries, >99.5%) were employed. All chemicals were used as received.

BaTiO3:Rh was prepared by a hydrothermal method as follows. The starting materials, Ba(OH)₂·8H₂O (Kanto Chemicals, >98.0%), TiO2, a rhodium source and 50 mL water, were placed in a 100 mL Teflon-lined stainless autoclave. The ratio of Ba/Ti/Rh was 1.0-1.1/0.98/0.02. The autoclave was then heated 433 K for 42 h. After the hydrothermal treatment, the products were naturally cooled down to room temperature. The resulting yellow precipitate was washed twice with hot water and four times with water, followed by drying at 363 K overnight. The powder was heated with or without addition of 5.0 mol% of $Ba(OH)_2 \cdot 8H_2O$ at 873-1423 K for 7 h.

4.2. Characterization of photocatalysts

The prepared samples were studied by powder X-ray diffraction (XRD; MiniFlex 600 Rigaku; Cu Kα), UV-visible diffuse reflectance spectroscopy (DRS; V-565, JASCO), scanning electron microscopy (SEM; JSM-7500F, JEOL), transmission electron microscopy (TEM; H-7650, Hitachi High-Technologies), and Xray photoelectron spectroscopy (XPS; ESCA-3400, Shimadzu). The binding energies determined by XPS were corrected in reference to the C1s peak (285.0 eV) for each sample. Elemental analysis of BaTiO₃:Rh was conducted using inductively coupled plasma mass spectrometry (ICP-MS; 7700x,

Technologies). The Brunauer-Emmett-Teller (BET) surface area was measured using a BELSORP-mini (BEL Japan) at liquid nitrogen temperature (77 K).

4.3. Photocatalytic reactions

The procedure was essentially the same as reported previously.6 Briefly, reactions were conducted at room temperature using a top-irradiation type cell that was connected to a closed gas circulation system made of glass. 100 mg of the as-prepared BaTiO3:Rh powder was dispersed in an aqueous methanol solution (100 mL, 10 vol%). A Pt cocatalyst was loaded onto BaTiO₃:Rh by an in situ photodeposition method using H₂- $PtCl_6 \cdot 6H_2O$ (Kanto Chemicals, >98.5%) as the precursor. After outgassing the reactant solution by a vacuum pump, Ar gas (ca. 5 kPa) was introduced into the reaction system and the solution was irradiated under a 300 W Xe lamp (Cermax, PE300BF) with an output current of 20 A. The irradiation wavelength was collected by the combination of a cold mirror (CM-1), cutoff filter (L42) and water filter ($\lambda > 420$ nm). The evolved gases were analyzed by gas chromatography (Shimadzu, GC-8A with TCD detector and MS-5A column, argon carrier gas). Apparent quantum yields (AQYs) for H2 evolution were measured in the same manner, as reported previously.6

Acknowledgements

The authors thank Dr Miharu Eguchi (National Institute of Materials Science, Japan) for assistance in TEM observation. This work was supported by the PRESTO/JST program "Chemical Conversion of Light Energy" and a Grant-in-Aid for Young Scientists (A) (Project 25709078).

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