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Flux-mediated doping of SrTiO₃ photocatalysts for efficient overall water splitting†

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SrTiO₃ is a photocatalyst that is well known for its activity for the overall water splitting reaction under UV light irradiation. In this study, the effects of SrCl₂ flux treatments and Al doping on the photocatalytic properties of SrTiO₃ were investigated. The SrTiO₃, which showed an apparent quantum efficiency of 30% at 360 nm in the overall water splitting reaction, the highest value reported so far, was prepared by SrCl₂ flux treatments in alumina crucibles. Scanning electron microscopy and X-ray diffractometry revealed that the flux-treated SrTiO₃ consisted of well-crystalline particles with a cubic shape reflecting the perovskite-type structure. Inductively coupled plasma optical emission spectroscopy revealed that Al ions from the alumina crucibles were incorporated into the SrTiO₃ samples. The SrTiO₃ that was treated with SrCl₂ flux in Al-free conditions showed a marginal improvement in photocatalytic activity despite the high crystallinity and the clear crystal habit. Doping SrTiO₃ with Al improved the photocatalytic activity even without SrCl₂ treatment. These results suggested that Al doping was a principal factor in the dramatic improvement in the water splitting activity of the flux-treated SrTiO₃. The effects of flux treatments and Al doping on the morphology and water splitting activity of SrTiO₃ were discussed separately.

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1 Introduction

Photocatalytic water splitting is a promising technology for the utilization of solar energy in the production of renewable hydrogen.^{1–4} In the last four decades, a number of particulate photocatalysts capable of overall water splitting have been developed.¹ La-doped NaTaO₃ loaded with NiO has reportedly split water at an apparent quantum efficiency of 56% at 270 nm.⁵ Zn-doped Ga₂O₃ loaded with Rh_{2–y}Cr_yO₃ has also been reported to split water at a high reaction rate.⁶ Some layered oxides such as K₄Nb₆O₁₇,⁷ Rb₄Nb₆O₁₇,⁸ and Rb₂La₂Ti₃O₁₀ (ref. 9) are known to show relatively high apparent quantum efficiencies (5%, 10%, and 5%, respectively) at 330 nm when modified with appropriate cocatalysts. However, these materials are

active only under UV light, limiting their application to solar energy conversion. The highest apparent quantum efficiency recorded in the visible light region was 5.1% (at 410 nm), attained by Rh_{2–y}Cr_yO₃/(Ga_{1–x}Zn_x)(N_{1–x}O_x).³ It is necessary to develop effective methods to improve the apparent quantum efficiency at wavelengths to achieve a solar-to-hydrogen energy conversion efficiency of 5–10%, which has been targeted for cost-competitive production of solar hydrogen *via* photocatalytic water splitting.^{3,4,10}

Given a specific photocatalytic material, controlling charge separation and migration is critical because the band structure and crystallographic character of each semiconducting material are unique. The movement of excited electrons and holes, and thus the photocatalytic activity, can be greatly affected by the crystallinity, particle size, and doping. Higher crystallinity and smaller particle sizes are desirable for charge migration toward surface active sites before recombination. Highly crystalline particles can be obtained by using a flux as a growth medium. The flux method, which allows the growth of crystalline particles *via* dissolution and recrystallization of solutes driven by supersaturation, has been applied to the synthesis of metal oxides,¹¹ including semiconducting oxides, such as K₄Nb₆O₁₇,¹² KNb₃O₈,¹³ Na₂Ti₆O₁₃,¹⁴ K₂Ti₆O₁₃,¹⁵ and SnNb₂O₆.¹⁶ Some of these oxides showed improved photocatalytic activity compared to those prepared by other synthesis methods. Some semiconducting (oxy)nitride photocatalysts such as LaTiO₂N,¹⁷ C₃N₄,¹⁸ and Ta₃N₅ (ref. 19) have also been prepared with the aid of a

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† Electronic supplementary information (ESI) available: DR spectra, XRD patterns, SEM images, and photocatalytic activity of the samples. See DOI: 10.1039/c5ta04843e

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flux. However, the downside of this method is the incorporation of impurities into the target material, a commonly observed phenomenon during flux treatment.²⁰ On the other hand, doping can often change the particle morphology dramatically⁵ and create mid-gap states essential for visible light activity of some wide-band-gap oxides.²¹ In fact, the high activity of NaTaO₃ and the visible light activity of SrTiO₃ photocatalysts rely on such doping effects. Recently, much effort has been made to understand the effect of doping in terms of carrier dynamics.^{22–24} From the material synthesis standpoint, the challenges in the activation of photocatalysts lie in how to lower defect densities, reduce particle sizes, and incorporate dopants effectively. The flux treatment of photocatalytic materials may offer a solution to such challenges.

SrTiO₃ is a classic photocatalyst that has been reported to be active in overall water splitting under UV light since 1980 (ref. 25 and 26) and is still widely investigated in fundamental studies on the effects of doping,^{21,23,27,28} particle morphology,²⁹ and cocatalysts.³⁰ In a recent study, we found that doping of lower-valence cations in SrTiO₃, such as Na⁺ into Sr²⁺ and Ga³⁺ into Ti⁴⁺, dramatically enhanced the photocatalytic activity during the overall water splitting reaction.²⁷ We attributed the positive effect of doping to the lower density of trivalent Ti states. Thus, the effects of the incorporation of even a small amount of impurity into SrTiO₃ during the flux treatment should be carefully investigated.

In this paper, we studied the effects of SrCl₂ flux treatment and doping on the physical properties and photocatalytic activity of SrTiO₃. The crystallinity and water splitting activity of SrTiO₃ were dramatically improved by the flux treatment in alumina crucibles. A small amount of Al doped into SrTiO₃ from an alumina crucible was found to be responsible for the enhancement in photocatalytic activity. It was found that the external doping of Al in the presence of flux was the most effective for controlled Al doping and produced an apparent quantum efficiency exceeding 30% at 360 nm, the highest value reported so far in this wavelength region and on SrTiO₃ powder.

2 Experimental

Sample preparation

As-purchased SrTiO₃. SrTiO₃ (Wako Pure Chemicals Industries, Ltd, 99.9%) was employed as a raw material without any post-treatment (hereafter STO(pristine)).

Flux-treated SrTiO₃. SrTiO₃ (Wako Pure Chemicals Industries, Ltd, 99.9%), Al₂O₃ (Sigma-Aldrich Co, LLC., nanopowder), and SrCl₂ (Kanto Chemicals Co., Inc., 98.0%, anhydrous) were used as raw materials. SrTiO₃ and SrCl₂ were thoroughly mixed in an agate mortar. The mixture was heated either in a yttria or an alumina crucible at 1100 °C for 10 h. After the mixture was cooled to room temperature, SrTiO₃ was separated from the solidified mass by repeated washing with deionized water until no white AgCl precipitate formed in rinse solutions upon adding AgNO₃. The SrTiO₃ samples treated in yttria and alumina crucibles will hereafter be referred to as STO(flux-Y) and STO(flux-Al), respectively, where -Y and -Al are used to highlight the material of the crucible used for the flux treatment. For STO(flux-

Al), the same procedure was followed, except that heating was performed at 900 and 1000 °C instead of 1100 °C. For the synthesis of STO(flux-Y) with Al dopant, Al₂O₃ was mixed together with SrTiO₃ and SrCl₂ and the same procedure was followed subsequently. The STO(flux-Y) samples with Al₂O₃ addition will hereafter be referred to as x%Al-STO(flux-Y), where x% represents the Al/Ti molar ratio in the starting mixture.

Al-doped SrTiO₃. SrTiO₃ (Wako Pure Chemicals Industries, Ltd, 99.9%) and Al₂O₃ (Sigma-Aldrich Co, LLC., nanopowder) were thoroughly mixed in an agate mortar. The mixture was heated in an alumina crucible at 1100 °C for 10 h. These samples will hereafter be referred to as Al-STO(ssr-Y).

Deposition of cocatalyst. A mixed oxide of rhodium and chromium, Rh_{2–y}Cr_yO₃, was loaded as a cocatalyst by the impregnation method. The details of the method used have been described elsewhere.³¹ The samples were loaded with Na₃RhCl₆·nH₂O (Mitsuwa Chemistry Co., Ltd, Rh 17.8 wt%) and Cr(NO₃)₃·9H₂O (Kanto Chemicals Co., Inc., 98.0–103.0%) as Rh and Cr sources, respectively, and calcined in air at 350 °C for 1 h.

Measurement of photocatalytic activity

The activities of the photocatalyst samples were tested in a closed gas circulation system with a top-irradiation-type reactor. The deionized water (100 mL) was evacuated to remove air completely. The reactor was irradiated using a 300 W xenon lamp ($\lambda > 300$ nm) through a quartz window or using a 450 W high-pressure mercury lamp through a quartz cooling jacket and, when necessary, a 2 cm-diameter slit, a band pass filter ($\lambda = 360$ nm, FWHM = 10 nm), and a series of neutral density filters (OD = 0.3, 0.5, 1.0, and 2.0) to irradiate the sample with monochromatic light with controlled intensity. The evolved gases were analyzed by a gas chromatograph (Shimadzu, GC-8A) equipped with a thermal conductivity detector, using Ar as a carrier gas.

Sample characterization

The crystal structures of the products were characterized by X-ray diffractometry (XRD; RINT Ultima III, Rigaku Co.) using Cu K α radiation at 40 kV and 40 mA. XRD peaks due to Cu K α ₁ and K α ₂ radiation were deconvoluted and the full width at half maximum (FWHM) of the (110) peak due to the Cu K α ₁ radiation was estimated. Specific surface areas were measured with a Belsorp-miniII (BEL Japan Inc.). The morphology of the powder was observed by scanning electron microscopy (SEM; S-4700, Hitachi High-Technologies Co.). Ultraviolet-visible diffuse reflectance spectrometry (DRS; V-670, Jasco Co.) was performed using spectralon (Jasco Co.) as a reference material. Inductively coupled plasma optical emission spectroscopy (ICP-OES; Shimadzu Co., ICPS-8100) was used for elemental analysis. SrTiO₃ powder (0.01 g) was melted with 1.0 g of a 3 : 1 mixture of Na₂CO₃ and B(OH)₃ by heating. An aqueous solution of tartaric acid (5%, 10 mL), HCl (1 + 1, 4 mL), and H₂O₂ (30 wt%, 1 mL) were added to dissolve the melt, and diluted with distilled water to make the total volume 100 mL. The resulting solution was used to measure Al and Y. The solution was further diluted tenfold to measure Sr and Ti.



3 Results and discussion

Commercially available SrTiO₃ (hereafter STO(pristine)) was mixed with SrCl₂ at a molar ratio of SrCl₂/SrTiO₃ = 10 and heated at 1100 °C for 10 h in yttria crucibles (hereafter STO(flux-Y)) and alumina crucibles (hereafter STO(flux-Al)) to obtain flux-treated SrTiO₃. X-ray diffraction (XRD) patterns are presented in Fig. 1. The XRD patterns for STO(flux-Y) and STO(flux-Al) showed only peaks attributable to the SrTiO₃ phase and exhibited sharper XRD peaks than STO(pristine), as reported in the literature.^{24,29} The FWHM of the (110) XRD peak was 0.103 for STO(pristine) and it was reduced to 0.071 for STO(flux-Y) (Table 1). Under the same measurement conditions, the difference in the FWHM of XRD peaks reflects the difference in size and strain of crystallites. The reduction of the FWHM after the flux treatment should reflect the growth of crystallites with weaker strain, which is indicative of higher crystallinity.

The SEM images of the samples in Fig. 2 show that STO(pristine) consisted of particles with irregular shapes, a few hundred nanometers in size. The flux treatment changed the

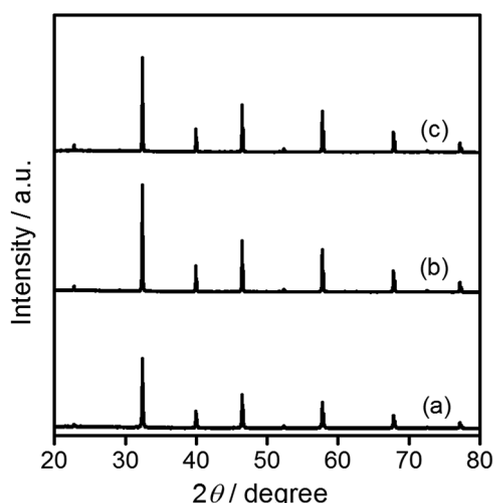


Fig. 1 XRD patterns for SrTiO₃ particles. (a) STO(pristine), (b) STO(flux-Y), and (c) STO(flux-Al).

Table 1 FWHM of the (110) diffraction peak, BET surface areas, and molar ratios, determined by ICP-OES of the SrTiO₃ photocatalysts

Sample	FWHM of (110) peak/°	BET surface area/m ² g ⁻¹	Molar ratio	
			2[Al]/([Sr] + [Ti])	2[Y]/([Sr] + [Ti])
STO(pristine)	0.103	3.6	0.04%	0.00%
STO(flux-Y)	0.071	1.5	0.02%	0.48%
STO(flux-Al), 1100 °C	0.067	0.9	0.31%	0.00%
STO(flux-Al), 1000 °C	0.074	1.3	0.11%	0.01%
STO(flux-Al), 900 °C	0.084	1.9	0.12%	0.00%

particle morphology dramatically. The STO(flux-Y) and STO(flux-Al) samples consisted of truncated cubic particles that exhibited a perovskite-type structure. The particles were 0.2–2 μm and 0.2–3 μm in size, respectively. The equilibrium crystal shape of SrTiO₃ was reported to be truncated cubic.³² Since the flux treated STO crystals also consisted of truncated cubes, it is considered that these crystals exposed the surfaces with the lowest formation energy. The morphological change agreed well with the change in specific surface area of the samples. The BET surface area decreased upon flux treatment (Table 1). The BET surface area for STO(flux-Al) was slightly smaller than that for STO(flux-Y), agreeing well with the difference in their particle sizes.

Diffuse reflectance spectra (DRS) of the SrTiO₃ samples showed that their band gap was not changed by the flux treatments (Fig. S1 in the ESI†). The absorption edge was located at approximately 390 nm. Their indirect band gaps,^{33,34} as estimated from the Tauc plots, were 3.2 eV, which well agreed with the reported values.³⁵

Fig. 3 shows the water splitting activities of STO(pristine), STO(flux-Y) and STO(flux-Al). The water splitting activity of STO(pristine) roughly tripled upon SrCl₂ flux treatment in an yttria crucible (STO(flux-Y)). This is presumably due to the improvement in crystallinity. When the same SrCl₂ flux treatment was carried out on SrTiO₃ in an alumina crucible (STO(flux-Al)), the water splitting activity was enhanced much more significantly, reaching 550 H₂ μmol h⁻¹ and 280 O₂ μmol h⁻¹. Our recent study on carrier dynamics in these SrTiO₃ powders conducted by time-resolved absorption (TA) spectroscopy revealed that most photoexcited electrons in STO(pristine) were deeply trapped, while those in STO(flux-Al) were in the conduction band or shallowly trapped.²⁴ This result well explains the enhanced water splitting activity of STO(flux-Al), because deeply trapped electrons would not be readily used for the hydrogen evolution reaction. STO(flux-Al) split water steadily for at least several hours, as shown in Fig. 4. The apparent quantum efficiency of STO(flux-Al) was measured to be 30% at 360 nm. This is much higher than the 4.3% reported for KCl-treated SrTiO₃, although the difference in reaction

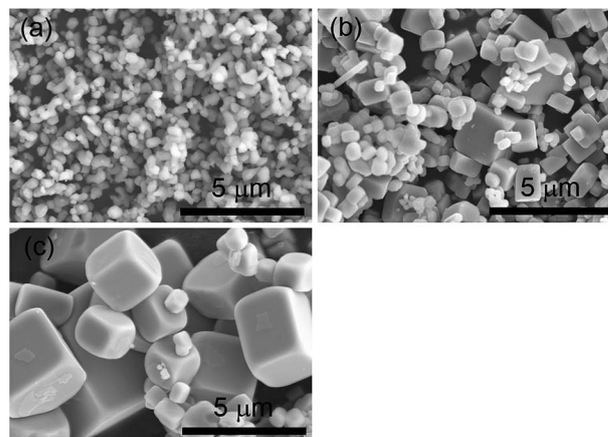


Fig. 2 SEM images of SrTiO₃ particles. (a) STO(pristine), (b) STO(flux-Y), and (c) STO(flux-Al).



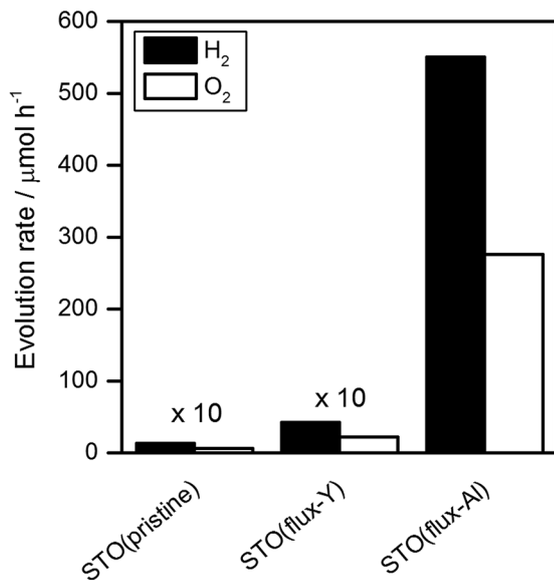


Fig. 3 Water splitting activity of SrTiO₃ photocatalysts. Reaction conditions: catalyst, 0.1 g; cocatalyst, Rh_{2-y}Cr_yO₃ (Rh 0.1 wt%, Cr 0.1 wt%); reaction solution, 100 mL H₂O; light source, 300 W Xe lamp ($\lambda > 300$ nm).

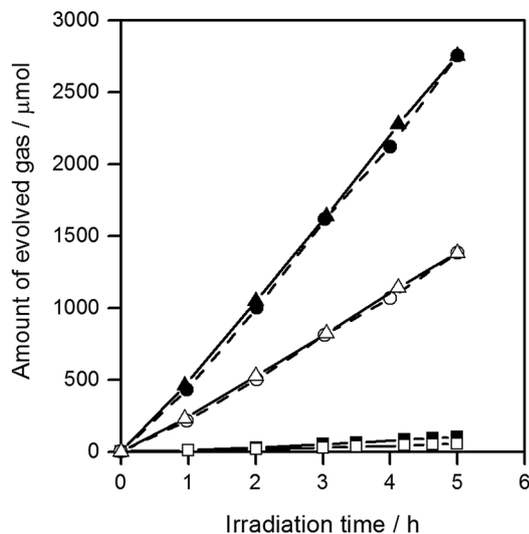


Fig. 4 Gas evolution during the water splitting reaction on STO(flux-Al). H₂ (■) and O₂ (□) on STO(flux-Al) treated at 900 °C, H₂ (●) and O₂ (○) on STO(flux-Al) treated at 1000 °C, and H₂ (▲) and O₂ (△) on STO(flux-Al) treated at 1100 °C. Reaction conditions: catalyst, 0.1 g; cocatalyst, Rh_{2-y}Cr_yO₃ (Rh 0.1 wt%, Cr 0.1 wt%); reaction solution, 100 mL H₂O; light source, 300 W Xe lamp ($\lambda > 300$ nm).

conditions should be taken into account.²⁹ We note that the apparent quantum efficiency was in fact dependent on the intensity of the incident light. As shown in Fig. S2 in the ESI,[†] the apparent quantum efficiency increased with light intensity under the experimental conditions examined. This result is not consistent with reaction orders to light intensity commonly observed in photocatalytic reactions, which is a first order.³⁶ We suspect that a certain kind of trap state with a limited density

may have to be filled with photoexcited carriers to attain sufficient photoconductivity for charge separation. The kinetic interpretation of the light intensity dependence of the photocatalytic water splitting rate will be reported in follow-up studies.

The FWHM of the (110) XRD peak for STO(flux-Al) treated at 1100 °C and STO(flux-Y) were 0.067 and 0.071, respectively (Table 1). Therefore, the crystallinity of STO(flux-Al) treated at 1100 °C was higher than that of STO(flux-Y). This might explain the large difference in the enhancement of photocatalytic activity by SrCl₂ flux treatment in alumina and yttria crucibles. To determine the influence of crystallinity on the photocatalytic activity of flux-treated SrTiO₃, the crystallinity of STO(flux-Al) was controlled by increasing the treatment temperature above the melting point of SrCl₂ (874 °C). For treatment temperatures of 900–1100 °C, the XRD patterns for all samples were attributed to single-phase SrTiO₃, regardless of the treatment temperature (Fig. S3 in the ESI[†]). However, the FWHM of the (110) XRD peaks of STO(flux-Al) increased upon lowering the flux treatment temperature and became comparable to that of STO(flux-Y) (Table 1). The water splitting activity of STO(flux-Al) became markedly higher when the treatment temperature increased from 900 to 1000 °C (Fig. 4). This suggested that the improvement in water splitting activity of flux-treated SrTiO₃ was associated with improved crystallinity. However, STO(flux-Al) treated at 900 and 1000 °C exhibited a significantly higher water splitting rate than STO(flux-Y), although the crystallinity of the former was inferior. This points to a different, more predominant factor governing the water splitting activity of SrTiO₃.

We speculated that Al impurity, derived from the alumina crucibles, became incorporated into SrTiO₃ during flux treatment. Indeed, it has been reported that doping SrTiO₃ with lower-valence cations can boost its photocatalytic activity.²⁷ Therefore, an elemental analysis of the samples was carried out by inductively coupled plasma optical emission spectroscopy (ICP-OES). Table 1 tabulates the compositions of flux-treated SrTiO₃. Al ions, derived from the alumina crucibles, were indeed detected in the STO(flux-Al) samples, where they would replace the Ti⁴⁺ sites of SrTiO₃. As expected, STO(flux-Y) was free of Al, but instead, it was doped with Y from the yttria crucibles. The higher water splitting activity of STO(flux-Al) compared to STO(flux-Y) is thus most likely attributable to the effect of Al doping. The enhancement of the water splitting activity by doping lower valence cations is consistent with the recent studies^{6,27,37} but contradictory to an earlier work on platinumized doped TiO₂ systems.³⁸ The difference probably resulted from the cocatalysts used. The Rh_{2-y}Cr_yO₃ cocatalyst is known to block the reverse reaction, *i.e.*, formation of water from hydrogen and oxygen molecules, unlike noble metals which readily catalyze the reverse reaction.^{34,39} Therefore, in this work, the water splitting activity is not obscured by the reverse reactions and closely reflects the change in the behavior of photoexcited charge carriers. Y doping did not have a significant positive effect on the photocatalytic activity. Unlike Al, Y presumably did not substitute well at the Ti sites because of the larger mismatch in ionic radius.

In an attempt to control Al doping in the presence of the SrCl₂ flux, SrTiO₃ and Al₂O₃ were mixed at molar ratios of



Al/Ti = 0.1%, 1%, and 10%, and heated together with the SrCl₂ flux in yttria crucibles. The resulting samples are referred to as x%Al-STO(flux-Y). The amount of Al doping was not directly proportional to the amount of Al₂O₃ added, but did increase with it (Table 2). In addition, certain amounts of Y were introduced from the yttria crucibles, similar to the case for the STO(flux-Y) sample. However, it is unlikely that Y doping at this level overwhelms the enhancement of photocatalytic activity by Al doping, considering the results for STO(flux-Y) (Fig. 3). The XRD patterns, SEM images, and BET surface areas for the samples and their water splitting activities are presented in Fig. S4 and S5 in the ESI,[†] Table 2, and Fig. 5, respectively. The 0.1%Al-STO(flux-Y) sample exhibited XRD patterns similar to those for STO(flux-Al). Single-phase SrTiO₃ was observed, and the FWHM of the (110) diffraction peak was 0.067. The morphology and BET surface area for 0.1%Al-STO(flux-Y) were also comparable to those for STO(flux-Al). However, the water

splitting activity of 0.1%Al-STO(flux-Y) was much lower than that of STO(flux-Al) due to the lower Al content in 0.1%Al-STO(flux-Y) compared to STO(flux-Al). The particle size of 1%Al-STO(flux-Y) and 10%Al-STO(flux-Y) decreased upon Al addition and most of the particles lost their crystal facets, which were distinctive for STO(flux-Al) and STO(flux-Y). This change in morphology is presumably due to suppression of the crystal growth of SrTiO₃ by the excess Al₂O₃, similar to the case for NaTaO₃ doped with La.⁵ A high activity for the overall water splitting reaction, comparable to that for STO(flux-Al), was obtained using yttria crucible when more than 1% of Al was added. This result supports our speculation that Al doping is the controlling factor for the enhancement of photocatalytic activity of SrTiO₃. On the other hand, the high activity of 10%Al-STO(flux-Y) despite its comparatively lower crystallinity may have resulted from the small particle sizes, which shorten the time needed for the migration of photoexcited carriers from the interior to the surface of photocatalyst particles.

To examine the effect of aluminum doping separately, Al₂O₃ was added as a dopant to SrTiO₃ at Al/Ti molar ratios ranging from 0.1% to 10%, and the mixtures were calcined in the absence of the SrCl₂ flux for a solid state reaction. The resulting products are referred to as x%Al-STO(ssr-Y), where x% represents the Al/Ti molar ratio in the starting mixture. As tabulated in Table 2, the Al content increased monotonically with increasing amount of Al₂O₃ addition, although the amounts detected were somehow lower than the amounts added to the starting material for high Al₂O₃ contents (Al >5%). The amount of Al incorporated into SrTiO₃ had a significant influence on the water splitting activity of the resulting samples. The water splitting activity peaked for 0.1%Al-STO(ssr-Y), as shown in Fig. 6. The activity of 0.1%Al-STO(ssr-Y) was two orders of magnitude higher than that of STO(pristine), but lower than that of STO(flux-Al) (Fig. 3). At this doping amount, no impurity

Table 2 FWHM of the (110) diffraction peak, BET surface areas, and molar ratios, determined by ICP-OES of the SrTiO₃ photocatalysts

Sample	FWHM of (110) peak/ ^o	BET surface area/m ² g ⁻¹	Molar ratio	
			2[Al]/([Sr] + [Ti])	2[Y]/([Sr] + [Ti])
0.1%Al-STO(flux-Y)	0.067	0.9	0.12%	0.16%
1%Al-STO(flux-Y)	0.096	2.7	1.01%	0.70%
10%Al-STO(flux-Y)	0.091	2.4	1.36%	0.57%
0.1%Al-STO(ssr-Y)	0.077	1.7	0.10%	0.00%
1%Al-STO(ssr-Y)	0.094	3.0	1.07%	0.00%
5%Al-STO(ssr-Y)	0.095	3.7	4.24%	0.01%
10%Al-STO(ssr-Y)	0.096	3.9	8.25%	0.01%

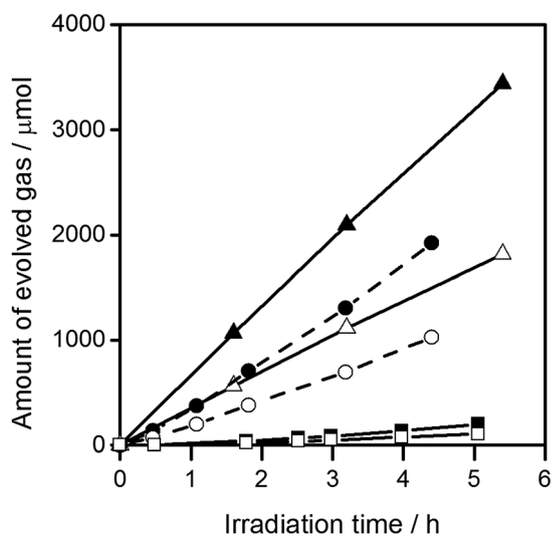


Fig. 5 Gas evolution during the water splitting reaction on Al-STO(-flux-Y). H₂ (■) and O₂ (□) on 0.1%Al-STO(flux-Y), H₂ (●) and O₂ (○) on 1%Al-STO(flux-Y), and H₂ (▲) and O₂ (△) on 10%Al-STO(flux-Y). Reaction conditions: catalyst, 0.1 g; cocatalyst, Rh_{2-y}Cr_yO₃ (Rh 0.1 wt%, Cr 0.1 wt%); reaction solution, 100 mL H₂O; light source, 300 W Xe lamp (λ > 300 nm).

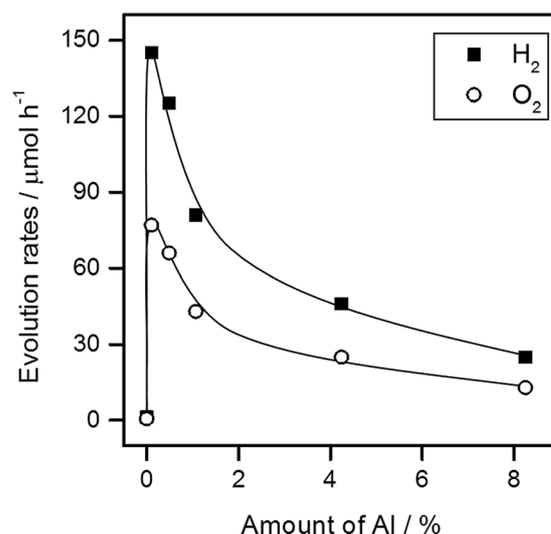


Fig. 6 Dependence of the water splitting activity of STO(ssr-Y) on the amount of Al doping. Reaction conditions: catalyst, 0.1 g; cocatalyst, Rh_{2-y}Cr_yO₃ (Rh 0.1 wt%, Cr 0.1 wt%); reaction solution, 100 mL H₂O; light source, 300 W Xe lamp (λ > 300 nm).



phase was detected in the XRD patterns of the sample (Fig. S6†). The BET surface area for 0.1%Al-STO(ssr-Y) decreased to $1.7 \text{ m}^2 \text{ g}^{-1}$ owing to the sintering process. However, it should be noted that no significant difference in morphology was observed between STO(pristine) and $x\%$ Al-STO(ssr-Y) (Fig. S7†). These results corroborate the necessity of the SrCl_2 flux for morphological change. It is worth mentioning that the impurity concentration that can dramatically improve the water splitting activity of SrTiO_3 ($\sim 0.1 \text{ mol}\%$) is close to the detection limits of commonly employed methods for elemental analysis, such as energy-dispersive X-ray emission spectroscopy (EDS) and X-ray photoelectron spectroscopy (XPS). Therefore, extra care needs to be taken when the effects of flux treatment on photocatalytic properties are discussed.

It should be noted that Al doping of SrTiO_3 was more effective when SrCl_2 was present during the heating. The doping amount of Al in the STO(flux-Al) sample from an alumina crucible may vary depending on the treatment conditions. Nevertheless, this sample showed a higher photocatalytic activity than the Al-STO(ssr-Y) samples containing various and controlled amounts of Al. It is thought that Al was not effectively doped into SrTiO_3 during the solid state reaction because Al had to diffuse from the outer surface of the particles. In contrast, a significant portion of the SrTiO_3 particles was once dissolved and recrystallized in the presence of SrCl_2 flux, together with alumina derived from the crucibles. During this process, some of the Al ions may be doped into the middle part of the SrTiO_3 particles and occupy the most stable state thermodynamically. As a result, Al doping can show stronger enhancement of photocatalytic activity when the SrCl_2 flux is used. Thus, it is concluded that the dramatic improvement in the photocatalytic activity of STO(flux-Al) was due to Al doping and the enhancement of crystallinity observed upon flux treatment.

4 Conclusions

The photocatalytic activity of SrTiO_3 in the overall water splitting reaction was dramatically improved by SrCl_2 flux treatment at $1100 \text{ }^\circ\text{C}$ in an alumina crucible. The improvement in activity was attributed mainly to the doping of Al derived from the crucibles. The morphological change and the enhanced crystallinity also improved the photocatalytic activity, although these factors were found to be less significant than the effect of Al doping on the basis of the results of flux treatment in Al-free conditions. It was confirmed that SrTiO_3 doped with Al under a SrCl_2 flux showed even higher water splitting activity than SrTiO_3 doped with Al by a solid state reaction. It is believed that the flux worked as a medium to dissolve the Al_2O_3 dopant and the host SrTiO_3 particles, facilitating the Al doping of SrTiO_3 . As a consequence, the apparent quantum efficiency in overall water splitting was increased to 30% at 360 nm. Flux-mediated doping is expected to greatly broaden the possibilities of photocatalytic materials by activating them under visible light irradiation. In addition, the incorporation of impurities into the samples during flux treatment was a common occurrence and could have strong impact on the photocatalytic activity.

Therefore, particular attention should be paid to flux treatment of photocatalysts.

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