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Effects of Rh-doping on the photooxidative degradation activity of titanate nanosheets†

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To investigate the Rh-doping effect on the photooxidative degradation activity of titanate nanosheets prepared by exfoliation of an $\text{H}_2\text{Ti}_3\text{O}_7$ crystal in aqueous media, a colloidal aqueous suspension of titanate nanosheets doped with Rh atoms at the Ti sites (TiNS:Rh_z , $z =$ amount of Rh, $[\text{Ti}_{3-z}\text{Rh}_z\text{O}_7]^{2-}$) was prepared. Oxidative degradation of methylene blue (MB) by TiNS:Rh_z in aqueous media under UV light irradiation was studied. Using X-ray diffraction, diffuse reflection spectroscopy, and photoelectrochemical measurements, the electronic band structure of TiNS:Rh_z was elucidated and was shown not to change with the amount of Rh doping at the Ti sites. However, the degradation of MB increased with an increase in Rh doping. From these results, it is suggested that the redox reaction between Rh^{3+} and Rh^{4+} strongly contributes to the oxidative degradation of MB in the presence of TiNS:Rh_z under UV light irradiation.

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1. Introduction

Two-dimensional titanate nanosheets (TiNSs), which can be prepared from layered titanate, $\text{H}_2\text{Ti}_n\text{O}_{2n+1}$ ($n \geq 3$), and/or lepidocrocite-type titanate, $\text{H}_n\text{Ti}_{2-n/4}\square_{n/4}\text{O}_4$ ($n \sim 0.7$; \square , vacancy), have attracted attention because they exhibit unique structural features such as high aspect ratios (thickness: 1 nm, surface area: \sim several μm^2) and malleability. Furthermore, they show useful physicochemical properties such as high cation-exchangeability, wide bandgap semiconductor properties, high dielectric constant, and photocatalytic ability.^{1–5} Thus, TiNSs have been studied by many researchers as a gas barrier, in water purification, as a functional coating reagent, as a self-cleaning material, and so on.^{6,7} However, the photocatalytic activity of these nanosheets is not sufficient for their application as a self-cleaning material. Thus, improvement in their photocatalytic activity is desired.

The metal doping method, in which other metal atoms are incorporated into the Ti sites of titanate, is a well-known method to improve photocatalytic activity. Improvement using this method originates from the formation of a new energy

band by orbital mixing and change in the carrier concentration.^{8,9} Adjustment of the photocatalytic properties of titanate affects its photoredox activity.

Recently, there have been many investigations into the metal doping of TiNSs. Ida *et al.* reported the photocatalytic activity of Rh-doped, lepidocrocite-type TiNSs, prepared by the exfoliation of $\text{H}_n\text{Ti}_{2-n/4}\square_{n/4}\text{O}_4$ ($n \sim 0.7$; \square , vacancy).¹⁰ In this work, it was reported that Rh atoms incorporated into the Ti sites of the TiNSs acted as reaction sites for photo-induced water splitting. Milanović *et al.* also reported the synthesis of Nb-doped, lepidocrocite-type TiNSs and the effect of Nb doping on the photodegradation of methylene blue (MB) in water purification.¹¹ Furthermore, Sasaki *et al.* reported the synthesis of lepidocrocite-type $\text{Ti}_{0.6}\text{Fe}_{0.4}\text{O}_2$ and $\text{Ti}_{0.8}\text{Co}_{0.2}\text{O}_2$ nanosheets.¹² It was shown that doping 3d metals into the Ti sites affected the crystal and electronic structures of the lepidocrocite-type TiNSs. The metal doping and effect of doping on the photocatalytic properties of TiNSs prepared by exfoliation of an $\text{H}_2\text{Ti}_n\text{O}_{2n+1}$ crystal have not been reported to date.

$\text{H}_2\text{Ti}_3\text{O}_7$ crystal can be prepared from layered $\text{Na}_2\text{Ti}_3\text{O}_7$ by the Na^+/H^+ ion-exchange. The basic layer architecture of $\text{H}_2\text{Ti}_3\text{O}_7$ was maintained nearly unchanged from that in $\text{Na}_2\text{Ti}_3\text{O}_7$ which the structure was determined by Andersson and Wadsley.¹³ $\text{Na}_2\text{Ti}_3\text{O}_7$ consists of edge-shared and corner-shared TiO_6 octahedral, which the sheet layers are separated by the Na^+ ion (as shown in Fig. S1 in ESI†). Unlike the lepidocrocite-type titanate, $\text{H}_n\text{Ti}_{2-n/4}\square_{n/4}\text{O}_4$ ($n \sim 0.7$; \square , vacancy), the $\text{H}_2\text{Ti}_3\text{O}_7$ crystal have no vacancies in their basic layer architecture. However, it is possible to dope some kinds of metals in Ti site such as Nb, Rh, Fe, Al, and *etc.* which their ionic radius not much different compared to Ti.

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† Electronic supplementary information (ESI) available: The schematic of $\text{H}_2\text{Ti}_3\text{O}_7$ crystal structure and their lattice parameters, the XRD patterns of Rh-doped sodium titanate, $\text{Na}_2\text{Ti}_{3-x}\text{Rh}_x\text{O}_7$, $\text{H}_2\text{Ti}_{3-x}\text{Rh}_x\text{O}_7$, and non-exfoliated TiNS:Rh_z , the AFM of TiNS:Rh_z , and XPS data for Ti and Rh compared between TiNS:Rh_0 and TiNS:Rh_{10} . See DOI: 10.1039/c6ra28731j



In this study, Rh-doped TiNSs (TiNS:Rh_z, *z* is the amount of Rh) were synthesised as a Na₂Ti_{3-x}Rh_xO₇ crystal by solid-state reaction. The electronic band structure of the synthesised TiNS:Rh_z was characterized by photocurrent and photoabsorption measurements. The photodegradation of MB in the presence of TiNS:Rh_z was investigated to evaluate its photocatalytic activity.

2. Experimental

2.1 Reagents and materials

Sodium carbonate Na₂CO₃ (99.9%) from Wako Pure Chemical Industries Co., anatase-type TiO₂ (99.7%) from High Purity Chemicals Co., Rh₂O₃ (99.9%) from Wako Pure Chemical Industries Co., methylamine (CH₃NH₂, 40 wt% solution) and tetramethylammonium hydroxide (TMAOH, 26 wt% solution) from Tokyo Chemical Industry Co., and MB (C₁₆H₁₈N₃-SCl·3H₂O, 3,7-bis(dimethylamino)-phenothiazin-5-ium chloride) from Kanto chemical Co. were used as received.

2.2 Synthesis of Rh-doped sodium titanate

Na₂Ti_{3-x}Rh_xO₇ was synthesized using a modified form of the solid-state reaction method reported by Izawa *et al.*¹⁴ as the following procedures. Na₂CO₃, anatase-type TiO₂ and Rh₂O₃ were mixed and grinded in a mortar. The mixture was calcined at 1173 K for 24 h in air. This operation was repeated again after grinding. The amount of Rh(*z*) added to the Ti sites was set from 0 to 10%.

2.3 Preparation of the nanosheet colloidal suspension

The colloidal suspension of TiNS:Rh_z was prepared according to a modified form of the method described by Miyamoto *et al.*¹⁵ 0.3 g of Na₂Ti_{3-x}Rh_xO₇ powder was dispersed in 30 cm³ of hydrochloric acid (1 mol dm⁻³) for exchanging the counter cation from Na⁺ to H⁺. Subsequently, it was shaken for 3 days at room temperature. The hydrochloric acid was replaced every day to allow the protonation reaction to proceed effectively. The filtrated H₂Ti_{3-x}Rh_xO₇ was washed with high purity water to remove any remaining hydrochloric acid, and was dried under a reduced pressure condition at room temperature overnight. The collected H₂Ti_{3-x}Rh_xO₇ was neutralized by methylamine aqueous solution at 333 K for 6 days to yield (CH₃NH₃)₂-Ti_{3-x}Rh_xO₇ powder. Subsequently, (CH₃NH₃)₂Ti_{3-x}Rh_xO₇ was dispersed in TMAOH solution ([Ti_{3-x}Rh_xO₇²⁻]/[TMAOH] = 5) by sonication for 5 days. Non-exfoliated TiNS:Rh_z was removed by centrifugation (at 4000 rpm or 1150g for 15 min, IEC61010-2-020, KUBOTA). The TiNS:Rh_z colloidal suspension was obtained as the supernatant.

2.4 Adsorption experiment

The adsorption experiment of MB¹⁶⁻¹⁸ by TiNS:Rh_z was carried out by the following procedure: (1) 0.01 cm³ of the TiNS:Rh_z (9.53 × 10⁻² mol dm⁻³, pH 11) was added to 10 cm³ of the MB aqueous solution (9.53 × 10⁻⁸ to 9.53 × 10⁻⁵ mol dm⁻³, pH 11), and this mixture was stirred at room temperature for 24 h, away from light. The molar ratio of the MB and TiNS:Rh_z was varied from 0.001 to 1.0. (2) After stirring, the mixture was filtrated

through a hydrophilic polyfluororesin membrane filter (pore size: 0.1 μm) to remove TiNS:Rh_z with MB from the suspension. (3) The amount of adsorbed MB by TiNS:Rh_z was calculated from the absorbance of MB in the filtrate at the peak wavelength of 663 nm. The extinction coefficient (ϵ) of the MB solution (pH 11) was 5.2 × 10⁴ dm³ mol⁻¹ cm⁻¹.

2.5 UV light irradiation experiment

To characterize the photocatalytic activity of TiNS:Rh_z in aqueous media, the solution of TiNS:Rh_z and MB was irradiated with UV light according to the following procedure: (1) the MB solution (4.7 × 10⁻⁶ mol) was mixed with TiNS:Rh_z (4.7 × 10⁻⁶ mol) to reach a total volume of 10 cm³ (pH 11), and stirred in the dark until the adsorption was at equilibrium. (2) UV light of 254 nm (0.88 mW cm⁻²) from a Xe lamp (MAX-301, Asahi Spectra Co.) was directed towards the mixture through a band pass filter, for a given time at room temperature. The irradiation area was confined to a square of 25 cm². (3) After the UV irradiation, the mixture was filtrated by the hydrophilic polyfluororesin membrane filter (pore size: 0.1 μm) to remove TiNS:Rh_z from the suspension. (4) The amount of adsorbed MB by TiNS:Rh_z was calculated from the absorbance of MB in the filtrate at the peak wavelength of 663 nm.

2.6 Characterization

X-ray diffraction (XRD) patterns were obtained by an X-ray diffractometer (RIGAKU, MiniFlex II) with Ni-filtered CuK α radiation (30 kV, 15 mA). The amount of Rh in TiNS:Rh_z was determined by inductively coupled plasma-atomic emission spectroscopy (ICP-AES: Optima -2000, Perkin-Elmer) after fully dissolving the prepared TiNS:Rh_z in nitric acid (1.0 mol dm⁻³). Absorption spectra of various samples were measured by a UV-Vis spectrophotometer (V-670, JASCO). Diffuse reflection spectra of the H₂Ti_{3-x}Rh_xO₇ powders were measured using the UV-Vis spectrophotometer (V-670, JASCO) attached to an integrating sphere system (ISN-723, JASCO).

According to the measurement setup reported by Usami *et al.*,^{19,20} photocurrent was measured by a three electrode electrochemical analysing system (Hokuto Denko, HSV-110) at a sweep rate of 10 mV s⁻¹. The TiNS:Rh_z was cast on FTO glass and dried at 333 K overnight. The TiNS:Rh_z film deposited on FTO glass acted as the working electrode, which was placed on a circle (9 mm in diameter) in a PTFE cell to adjust the electrode area to 0.636 cm². An Ag/AgNO₃ reference electrode and Pt wire counter electrode were immersed in an acetonitrile solution of 0.1 mol dm⁻³ tetraethylammonium perchlorate. A parallel beam of excitation light from a 150 W Xe lamp (Hamamatsu Photonics, L2195) was focused on the semiconductor layer of the FTO electrode through a light chopper (NF Electric Instruments, 5584A). The modulated photocurrent at the applied potential was amplified using a digital lock-in-amplifier (NF Electric Instruments, LI5640) and transferred to a PC to be plotted against the applied potential.





Fig. 1 The Tyndall effect observed by passing a laser beam through the TiNS:Rh_z colloidal suspension ($\sim 0.5 \text{ mmol dm}^{-3}$).

3. Results and discussion

3.1 Characterization of TiNS:Rh_z

Fig. 1 is a photograph of the prepared TiNS:Rh_z colloidal suspension, which is transparent but shows scattering of an incident laser beam because of the Tyndall effect. It was found that TiNS:Rh_z from $\text{Na}_2\text{Ti}_{3-x}\text{Rh}_x\text{O}_7$ could be exfoliated in all of the suspensions by the described procedures. The chemical compositions of the prepared TiNS:Rh_z are shown in Table 1. The actual amount of Rh in TiNS:Rh_z was lower than what was initially added. Some Rh may have been removed during protonation, intercalation of methylamine, or exfoliation by TMAOH (see Fig. S2 and S3 in ESI[†]). In the XRD patterns of the cast films prepared from the TiNS:Rh_z colloidal suspension, diffraction peaks originating from the deposited unknown crystal compounds were not found, since the deposited unknown crystal compounds were not layered materials and could not be exfoliated by the TMAOH aqueous solution.

Fig. 3 shows the UV-visible absorption spectra of all the TiNS:Rh_z colloidal suspensions of the same concentration $1 \text{ (} 2 \text{ mg dm}^{-3}\text{)}$. The band edge originates from the band gap of the TiNS:Rh_z, which exhibits a blue shift with respect to the diffuse reflectance spectra of the $\text{H}_2\text{Ti}_3\text{O}_7$ powder. This blue shift can be explained by the quantum-size effect.^{21,22} The TiNS:Rh_z existed as nanosheets with a thickness about 1.5 nm in the colloidal suspension (see Fig. S4 in ESI[†]). The optical band gap (E_g , estimated from the UV-vis absorption spectra of the TiNS:Rh_z colloidal suspension) and the conduction band level (E_{CB} , estimated from the photocurrent measurement of the TiNS:Rh_z cast film) are shown in Table 1. Both E_g and E_{CB} of all samples were not influenced by Rh doping into the Ti sites of the lattice.

3.2 The adsorption of methylene blue molecules by TiNS:Rh_z

The adsorption of MB by TiNS:Rh_z was evaluated by plotting the molar amount of adsorbed MB per $[\text{Ti}_{3-x}\text{Rh}_x\text{O}_7^{2-}]$ unit, C^{abs}

Table 1 The chemical formula, optical band gap (E_g), and conduction band potential (E_{CB}) of the TiNS:Rh_z samples

Sample	Chemical formula	E_g (eV)	E_{CB} (V vs. NHE)
TiNS:Rh0	$[\text{Ti}_3\text{O}_7]^{2-}$	4.20 ± 0.04	0.00
TiNS:Rh1	$[\text{Ti}_{2.999}\text{Rh}_{0.001}\text{O}_7]^{(2+\delta)-}$	4.17 ± 0.04	0.00
TiNS:Rh2	$[\text{Ti}_{2.992}\text{Rh}_{0.008}\text{O}_7]^{(2+\delta)-}$	4.17 ± 0.04	0.00
TiNS:Rh5	$[\text{Ti}_{2.970}\text{Rh}_{0.030}\text{O}_7]^{(2+\delta)-}$	4.15 ± 0.04	-0.01
TiNS:Rh10	$[\text{Ti}_{2.952}\text{Rh}_{0.048}\text{O}_7]^{(2+\delta)-}$	4.18 ± 0.04	-0.01

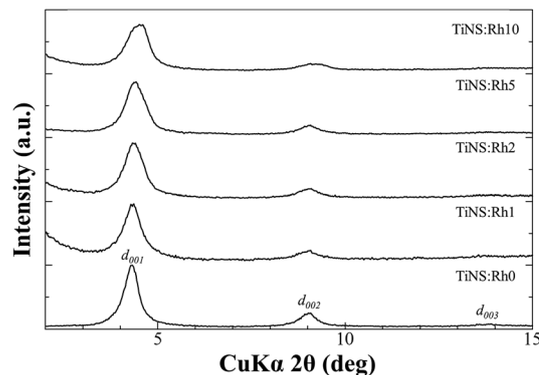


Fig. 2 XRD patterns of TiNS:Rh_z films cast on quartz substrates.

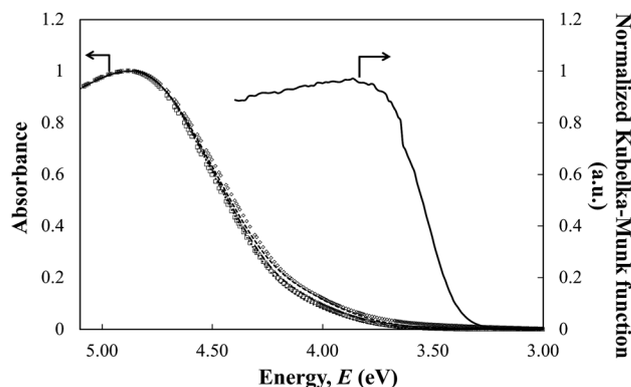


Fig. 3 The UV-vis absorption spectra of the TiNS:Rh_z colloidal suspension (concentration of 2 mg dm^{-3}) and diffuse reflectance spectra of the $\text{H}_2\text{Ti}_3\text{O}_7$ powders. ((●) TiNS:Rh0, (Δ) TiNS:Rh1, (—●) TiNS:Rh2, (◇) TiNS:Rh5, (□) TiNS:Rh10, and (—) $\text{H}_2\text{Ti}_3\text{O}_7$).

(mol mol^{-1}) against the molar amount of added MB per $[\text{Ti}_{3-x}\text{Rh}_x\text{O}_7^{2-}]$ unit, C^0 (mol mol^{-1}), as shown in Fig. 4. The value of C^{abs} increased along with an increase in the C^0 value. Therefore, the amount of adsorbed MB at saturation point could be calculated, and this is shown in Table 2. The adsorbed

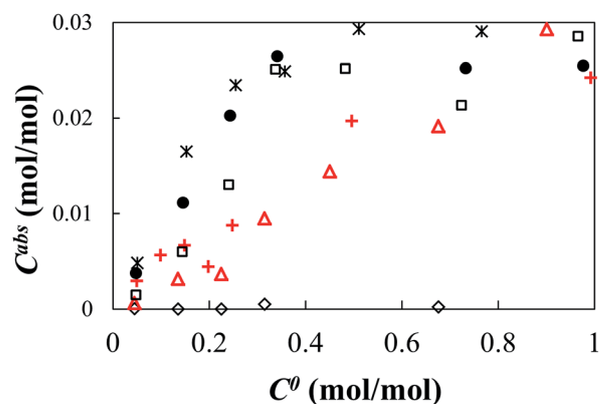


Fig. 4 The adsorption isothermal curves of MB by TiNS:Rh_z. ((◇) Absence of TiNS:Rh_z, (●) TiNS:Rh0, (Δ) TiNS:Rh1, (□) TiNS:Rh2, (×) TiNS:Rh5, and (●) TiNS:Rh10).



Table 2 Amount of absorbed MB by TiNS:Rh_z

Sample	Saturated amount of absorbed MB per molar amount of TiNS:Rh _z at equilibrium (mg g ⁻¹)
TiNS:Rh0	37 ± 3
TiNS:Rh1	36 ± 3
TiNS:Rh2	35 ± 3
TiNS:Rh5	31 ± 3
TiNS:Rh10	29 ± 3

amount was much lower than the formal anionic charge of TiNS:Rh_z (2 mol mol⁻¹) in all cases because of the competitive adsorption of MB and TMAOH. Table 2 also shows that the amount of adsorbed MB slightly decreased with an increase in Rh doping. This can be explained by the anionic charge of TiNS:Rh_z. In the case of TiNS:Rh_z ($z > 0$), the anionic charge changed because the ionic state of Rh was Rh³⁺ or Rh⁴⁺. This led to an increase in the anionic charge of TiNS:Rh_z. However, the result did not affect the MB adsorption because it strongly interacted with the TMA cation. As a result, the adsorption of MB decreased when Rh was incorporated into the Ti sites.

The shape of the adsorption isothermal curve was different for each sample. The C^{abs} value of the TiNS:Rh0 and TiNS:Rh1 linearly increased with an increase in the C^0 value. The adsorption isothermal curve of TiNS:Rh1 behaved similarly to TiNS:Rh0 but the C^{abs} value of TiNS:Rh2, TiNS:Rh5 and TiNS:Rh10 logarithmically increased. Thus, these results suggest that the surface properties of TiNS varied with Rh doping into the Ti sites.

3.3 Photodegradation of methylene blue

Degradation of MB by UV light irradiation was observed by changes in its UV-visible absorption spectra. Fig. 5 shows the UV-visible absorption spectra of filtrated MB solution from the mixture with TiNS:Rh0 after irradiation by UV light with a wavelength of 254 nm, at different irradiation times. The spectra of the solution were similar with a gradual decrease in absorbance seen for the peak at around 663 nm, originating from MB. This was observed for all TiNS:Rh_z samples, which

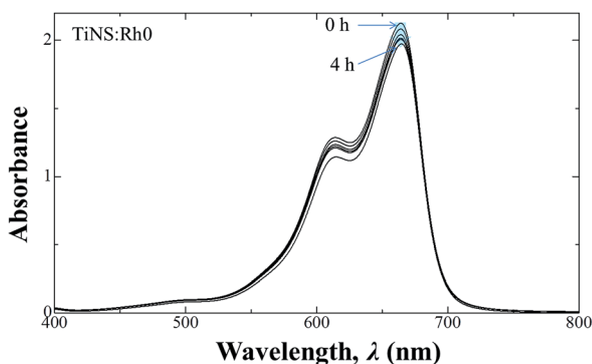


Fig. 5 UV-vis absorption spectral changes of MB solution under UV light (at a wavelength of 254 nm) irradiation in the presence of TiNS:Rh0.

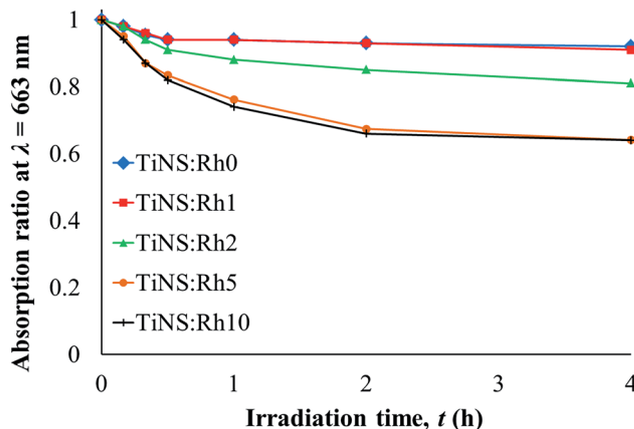


Fig. 6 The photodegradation ratio curves of the maximum absorption band of MB in the presence of the different TiNS:Rh_z samples (at 663 nm), after the UV light ($\lambda = 254$ nm) irradiation over different irradiation times.

shows that the MB was degraded by UV light irradiation and that TiNS:Rh_z acted as a photocatalyst to decompose MB.

To compare photocatalytic activity of each TiNS:Rh_z series, the absorbance ratio values at 663 nm (Abs_t/Abs_0 ; where Abs_t is an absorbance at each interval and Abs_0 is the initial absorbance or absorbance at $t = 0$) are plotted against the irradiation time in Fig. 6. All TiNS:Rh_z samples photocatalytically degraded MB. However, the duration of the reaction varied. TiNS:Rh0 and TiNS:Rh1 showed almost the same photodegradation ratio curve, while TiNS:Rh2, TiNS:Rh5, and TiNS:Rh10 produced a different result. This indicates that photocatalytic activity can be improved by Rh doping at the Ti sites of TiNS:Rh_z.

The quantum yield (Φ_{deg}) of the photodegradation reaction of MB by TiNS:Rh_z at initial time was calculated by:

$$\Phi_{deg} = \frac{\text{number of degraded MB molecules}}{\text{number of incident photons}}$$

Table 3 shows the Φ_{deg} value of each TiNS:Rh_z sample. TiNS:Rh0 and TiNS:Rh1 exhibit almost the same Φ_{deg} value. In contrast, the Φ_{deg} value of TiNS:Rh2, TiNS:Rh5, and TiNS:Rh10 are higher than that of TiNS:Rh0; thus, Φ_{deg} increased with increasing Rh doping. A similar divide between samples was observed in the adsorption study (see Table 2). Thus, it is evident that Rh doping modifies the surface properties of TiNS:Rh_z. However, the adsorption ability was not improved with an increase in Rh doping. This may be because the size of TiNS:Rh_z decreases with increase in Rh doping (see Fig. 2).

Table 3 Quantum yield (Φ_{deg}) of MB photodegradation

Sample	Φ_{deg} (%)
TiNS:Rh0	0.30
TiNS:Rh1	0.31
TiNS:Rh2	0.34
TiNS:Rh5	0.81
TiNS:Rh10	0.97



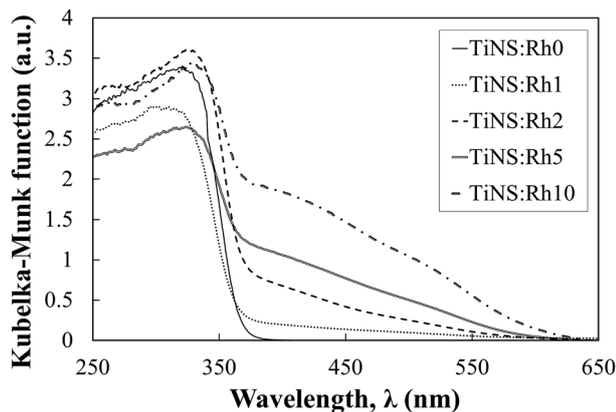


Fig. 7 The diffuse reflection spectra of the $\text{H}_2\text{Ti}_{3-x}\text{Rh}_x\text{O}_7$ powders.

Moreover, the difference in photocatalytic ability cannot be explained by the energy band structure because the band gap energy and band edge potentials of the conduction and valence bands did not vary with Rh doping (see Table 1).

Fig. 7 shows the diffuse reflection spectra of the $\text{H}_2\text{Ti}_{3-x}\text{Rh}_x\text{O}_7$ powders in terms of the Kubelka–Munk function. The absorption band edge at 350 nm originating from the band gap transition is observed in all of the TiNS:Rh $_z$ spectra. A broad absorption band appeared in the wavelength range from 350 to 600 nm when Rh atoms were incorporated into the Ti sites. The intensity of these new bands increased with an increase in Rh doping. According to previous results reported by Kudo *et al.*, Rh $^{3+}$ and Rh $^{4+}$ incorporated into the Ti sites of SrTiO $_3$ showed broad absorption bands around a wavelength of 400 and 500 nm, respectively.^{23–27} Hence, Rh dopants in the present work existed as both Rh $^{3+}$ and Rh $^{4+}$. This also has been confirmed by measuring the X-ray photoelectron spectroscopy. As a result, the peak of Rh 3d at 308.9 eV and 309.9 eV could be found, and thus this result shows the existence of Rh $^{3+}$ and Rh $^{4+}$ (see also in Fig. S5 and S6 of ESI†). The different photocatalytic activities of TiNS:Rh $_z$ in MB photodegradation may be explained by the existence of both Rh $^{3+}$ and Rh $^{4+}$. Irradiation of TiNS:Rh0 by UV light is known to cause oxidative degradation of MB by photo-induced holes, OH radicals, and superoxide anions.^{28–31} The redox cycle of Rh $^{3+}$ and Rh $^{4+}$ in TiNS:Rh $_z$ ($z > 0$) can be driven by band gap excitation of TiNS under UV light irradiation due to a hole pool effect of Rh atoms in six fold coordination with the oxygen atoms.³² Thus, the improvement of photocatalytic activities of TiNS:Rh $_z$ can be caused by the redox cycle of the Rh cations (Rh $^{3+}$ and Rh $^{4+}$) under UV light irradiation. However, it is still unclear whether the redox cycle of the Rh cations in the studied TiNS:Rh $_z$ samples was driven by the UV light. This will be the next challenging topic for research not only in titanate nanosheets but also in other metal oxide nanosheets and oxide materials.

4. Conclusions

In this study, the synthesis and photoelectrochemical characterization of TiNS:Rh $_z$ was performed. The band structure of the

TiNS:Rh $_z$ was not affected by Rh incorporation into the Ti sites. However, both adsorption and photodegradation behaviour of MB by TiNS:Rh $_z$ varied with Rh doping. The photodegradation activity of TiNS:Rh $_z$ was significantly improved with increased Rh doping, suggesting that the redox cycle of Rh $^{3+}$ and Rh $^{4+}$ plays an important role in the photodegradation of MB under UV light irradiation in the presence of TiNS:Rh $_z$.

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References

- H. Sato, K. Ono, T. Sasaki and A. Yamagishi, *J. Phys. Chem. B*, 2003, **107**, 9824–9828.
- T. Sasaki, Y. Ebina, M. Watanabe and G. Decher, *Chem. Commun.*, 2000, 2163–2164.
- T. Sasaki, Y. Ebina, Y. Kitami, M. Watanabe and T. Oikawa, *J. Phys. Chem. B*, 2001, **105**, 6116–6121.
- T. Sasaki, Y. Ebina, K. Fukuda, T. Tanaka, M. Harada and M. Watanabe, *Chem. Mater.*, 2002, **14**, 3524–3530.
- T. Maluangnont, K. Matsuba, F. Geng, R. Ma, Y. Yamauchi and T. Sasaki, *Chem. Mater.*, 2013, **25**, 3137–3146.
- X. Dong, M. Osada, H. Ueda, Y. Ebina, Y. Kotani, K. Ono, S. Ueda, K. Kobayashi, K. Takada and T. Sasaki, *Chem. Mater.*, 2009, **21**, 4366–4373.
- M. Harada, T. Sasaki, Y. Ebina and M. Watanabe, *J. Photochem. Photobiol., A*, 2002, **148**, 273–276.
- T. Sasaki, Y. Ebina, Y. Kitami, M. Watanabe and T. Oikawa, *J. Phys. Chem. B*, 2001, **105**, 6116–6121.
- H. Sun, G. Zhou, S. Liu, H. M. Ang, M. O. Tadé and S. Wang, *Chem. Eng. J.*, 2013, **231**, 18–25.
- S. Ida, N. Kim, E. Ertekin, S. Takenaka and T. Ishihara, *J. Am. Chem. Soc.*, 2015, **137**, 239–244.
- M. Milanović, L. M. Nikolić, I. Stijepović, A. G. Kontos and K. P. Giannakopoulos, *Mater. Chem. Phys.*, 2014, **148**, 874–881.
- Y.-H. Kim, M. Osada, L. Dong, H.-J. Kim and T. Sasaki, *J. Ceram. Soc. Jpn.*, 2015, **123**, 335–339.
- S. Andersson and A. D. Wadsley, *Acta Crystallogr.*, 1961, **14**, 1245–1249.
- H. Izawa, S. Kikkawa and M. Koizumi, *J. Phys. Chem.*, 1982, **86**, 5023–5026.
- N. Miyamoto, K. Kuroda and M. Ogawa, *J. Mater. Chem.*, 2004, **14**, 165.
- W. Kuo and P. Ho, *Dyes Pigm.*, 2006, **71**, 212–217.
- W. Kuo and P. Ho, *Chemosphere*, 2001, **45**, 77–83.



- 18 M. Feng, W. You, Z. Wu, Q. Chen and H. Zhan, *ACS Appl. Mater. Interfaces*, 2013, **5**, 12654–12662.
- 19 H. Usami and S. Sasaki, *J. Photopolym. Sci. Technol.*, 2013, **26**, 207–211.
- 20 H. Usami, Y. Iijima, Y. Moriizumi, H. Fujimatsu, E. Suzuki and H. Inoue, *Res. Chem. Intermed.*, 2007, **33**, 101–110.
- 21 T. Sasaki, M. Watanabe, H. Hashizume, H. Yamada and H. Nakazawa, *J. Am. Chem. Soc.*, 1996, **118**, 8329.
- 22 M. Osada and T. Sasaki, *Adv. Mater.*, 2012, **24**, 209–228.
- 23 A. Kudo and Y. Miseki, *Chem. Soc. Rev.*, 2009, **38**, 253–278.
- 24 R. Konta, T. Ishii, H. Kato and A. Kudo, *J. Phys. Chem. B*, 2004, **108**, 8992–8995.
- 25 K. Iwashina and A. Kudo, *J. Am. Chem. Soc.*, 2011, **133**, 13272–13275.
- 26 S. Kawasaki, K. Nakatsuji, J. Yoshinobu, F. Komori, R. Takahashi, M. Lippmaa, K. Mase and A. Kudo, *Appl. Phys. Lett.*, 2012, **101**, 033910.
- 27 S. Kawasaki, K. Akagi, K. Nakatsuji, S. Yamamoto, I. Matsuda, Y. Harada, J. Yoshinobu, F. Komori, R. Takahashi, M. Lippmaa, C. Sakai, H. Niwa, M. Oshima, K. Iwashina and A. Kudo, *J. Phys. Chem. C*, 2012, **116**, 24445–24448.
- 28 A. Mills and J. Wang, *J. Photochem. Photobiol., A*, 1999, **127**, 123–134.
- 29 R. Matthews, *Water Res.*, 1991, **25**, 1169–1176.
- 30 A. Houas, *Appl. Catal., B*, 2001, **31**, 145–157.
- 31 K. Wetchakun, N. Wetchakun and S. Phanichphant, *Desalin. Water Treat.*, 2015, **57**, 10286–10294.
- 32 J. Kunczewicz and B. Ohtani, *Chem. Commun.*, 2015, **51**, 298–301.

