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Silver-loaded sodium titanate photocatalysts for selective reduction of carbon dioxide to carbon monoxide with water

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Abstract: To obtain more efficient photocatalyst for photocatalytic reduction of CO2 with H2O and figure out the reason for nonstoichiometric O2 evolution, silver-loaded sodium titanate photocatalysts were further studied in the improved reaction conditions. After preliminary tests for two kinds of sodium titanate samples with different ratio of sodium to titanium (Na2Ti6O13 and Na2Ti3O7), several sodium hexatitanate (Na2Ti6O13) photocatalysts were further prepared in the flux method by changing the parameters such as the flux, the loading amount of Ag cocatalyst, and the loading method of the Ag cocatalyst. As a result, a Ag/Na2Ti6O13 sample prepared in a sodium chloride flux, with 1.0 wt% of Ag cocatalyst loaded by a photodeposition method, exhibited the highest production rate (4.6 μmol h⁻¹) and the highest selectivity (74%) to carbon monoxide among the examined samples, which are more than 29 times higher production rate and 2.7 times higher selectivity to carbon monoxide than those in the previous report. Furthermore, although required oxygen production rate equivalent to the formation rates of the reduced products was not observed in the previous study, it was found that the developed Ag/Na2Ti6O13(NaCl) photocatalyst produced enough amount of oxygen after a long induction period of 50 h in the present condition. The reasons for the insufficient oxygen formation in the initial period were also investigated and clarified, i.e., the chloride residues and the photoadsorption of O2 on the surface are responsible for the insufficient O2 evolution at the initial period.

Keywords: Photocatalytic conversion of CO2; Carbon monoxide; Sodium titanate; Silver cocatalyst.

1. Introduction

Photocatalytic CO2 reduction with water using solar energy is the desirable method for not only decreasing CO2 emission, but also producing valuable chemicals that can be used directly as fuels or in further chemical syntheses [1–5]. Heterogeneous photocatalysis have been studied for the reduction of CO2 with water, and the products reported are formate, CO, CH3OH and CH4 [6–14]. Among these possible reductive products from CO2, CO is the most useful product since it is a valuable chemical intermediate for further chemical syntheses and easily-separable from the aqueous reaction media to the gas phase. To obtain CO as a product, Ag cocatalyst was discovered [15], which has been widely studied in recent years, such as Ag/BaLa4Ti4O15 [15], Ag/Ga2O3 [16–20], Ag/KCaSrTa5O15 [21], Ag/La2Ti2O7 [22], Ag/CaTiO3 [23,24], and Ag/SrNb2O6 [25,26]. However, the photocatalytic activity has not been enough for the practical application. Thus, it is still imperative to develop highly efficient photocatalysts for this reaction.

In these reaction systems, it can be assumed that two reactions competitively occur, i.e., photocatalytic reduction of CO2 to form CO and O2 (Eq. 1) and photocatalytic water splitting into H2 and O2 (Eq. 2). In details, the reduction
of CO₂ and protons with the photoexcited electrons proceed to form CO and H₂, respectively on the reductive sites of the photocatalyst surface (Eqs. 3 and 4), while the oxidation of water with positive holes into O₂ and protons occurs on the oxidative sites (Eq. 5).

\[
\begin{align*}
\text{CO}_2 & \rightarrow \text{CO} + \frac{1}{2} \text{O}_2 & (1) \\
\text{H}_2\text{O} & \rightarrow \text{H}_2 + \frac{1}{2} \text{O}_2 & (2) \\
\text{CO}_2 + 2 \text{H}^+ + 2 e^- & \rightarrow \text{CO} + \text{H}_2\text{O} & (3) \\
2 \text{H}^+ + 2 e^- & \rightarrow \text{H}_2 & (4) \\
2 \text{H}_2\text{O} + 4 h^+ & \rightarrow \text{O}_2 + 4 \text{H}^+ & (5)
\end{align*}
\]

When no other reductive products than CO and H₂ are observed in the reaction system, the selectivity toward CO evolution, \( S_{\text{CO}}(\%) \), and the ratio of the consumed electron and hole, \( R(e^-/h^+) \), can be calculated according to Eqs. 6 and 7, respectively [24,26], where the production rate of CO, H₂ and O₂ are referred to as \( R_{\text{CO}} \), \( R_{\text{H}_2} \) and \( R_{\text{O}_2} \), respectively. The value of \( R(e^-/h^+) \) should be unity if the two competitive reactions, Eqs. 1 and 2, take place ideally in any balance.

\[
\begin{align*}
S_{\text{CO}} (%) &= 100 \times \frac{R_{\text{CO}}}{R_{\text{CO}} + R_{\text{H}_2}} & (6) \\
R(e^-/h^+) &= \frac{(2R_{\text{CO}} + 2R_{\text{H}_2})}{4R_{\text{O}_2}} & (7)
\end{align*}
\]

In our previous study [27], we focused on sodium hexatitanate (Na₂Ti₆O₁₃) since titanium and sodium are abundant elements in the Earth’s crust. The sodium hexatitanate samples were prepared by a flux method, on which Ag cocatalyst was loaded. The Ag/Na₂Ti₆O₁₃ photocatalyst exhibited the photocatalytic activity for the CO₂ reduction to CO. The photocatalyst consisted of unique rod-like fine crystal particles showing facets, and the Ag cocatalyst nanoparticles were deposited on the selected facets, especially after the photocatalytic reaction test. In this system, however, two problems remained; one was the low activity of CO evolution with low selectivity, such as \( R_{\text{CO}}=0.16 \mu\text{mol h}^{-1} \) and \( S_{\text{CO}}=27\% \), and the other was the uncertainty of the oxidation reaction, i.e., insufficient oxygen evolution, \( R(e^-/h^+)<1 \). As for the former problem, the reaction conditions should be reconsidered since we found recently that the reaction conditions, such as light intensity and CO₂ concentration in the reaction mixture, much affect the production rates of carbon monoxide and oxygen [24]. As for the latter problem, so far, many photocatalytic systems have faced this problem, i.e., the ratio of the consumed electron and hole \( R(e^-/h^+) \) estimated from the production rates was not unity, which is not consistent to the stoichiometric ratio. Among the proposed explanations, the most convinced one would be that the difficulty of water oxidation is due to the high overpotential of O₂ evolution, which originates from charge trapping by surface states and slow oxygen evolution kinetics [28–30]. This is also an explanation why some photocatalytic systems for water splitting produce H₂ without sufficient amount of O₂ (or \( R_{\text{H}_2}/R_{\text{O}_2}>2 \)) [31]. In addition, photoadsorption of O₂ on the photocatalyst surface [32,33] and the competitive oxidation of water to hydrogen peroxide [34,35] are also possible in the photocatalytic reaction with water. What’s more, for some special reaction systems using catalyst samples containing chloride residues originated from the catalyst preparation, such as a flux method with chloride flux, the residues of chloride may compete with oxygen evolution. Some researchers suggested that the Cl⁻ that exists in the photocatalytic system can react with the photogenerated holes [36,37]. Li and co-workers claimed that the existence of Cl⁻ affected both the H₂ evolution and O₂ evolution [38]. Furthermore, Iguchi et al. also reported that hypochlorous acid (HClO) was produced as an oxidation product of Cl⁻ with the formation of reduction products such as CO and H₂ under photoirradiation [39].
Thus, in the present study, the silver-loaded sodium titanate photocatalysts, one of which was once reported in the previous study [27], were again examined in the improved reaction conditions for the photocatalytic reduction of carbon dioxide by water. Two points were claimed in this study. First, after testing the composition of sodium titanate (Na₂Ti₆O₁₃ and Na₂Ti₃O₇), the performance of the prepared Ag/Na₂Ti₆O₁₃ catalysts were much improved by optimizing several conditions such as the flux in a flux method, and the loading method and the loading amount of silver cocatalyst. Second, the reasons for the nonstoichiometric O₂ evolution in the initial period were clarified.

2. Experimental
2.1. Sample preparation

The Na₂Ti₃O₇ sample was prepared by a solid state reaction (SSR) method reported in the literature [39]. The precursors (Na₂CO₃ (Kishida): TiO₂ (rutile, Kojundo)=1:3) were mixed for 30 minutes in an alumina mortar and calcined at 1073 K for 20 h, followed by additional grinding at room temperature and heating at 1073 K for another 20 h. Although it was examined to prepare another Na₂Ti₃O₇ sample by a flux method, no sample showing XRD patterns was obtained.

For the preparation of another Na₂Ti₆O₁₃ sample in the SSR method, the precursors were mixed for 30 minutes in an alumina mortar, where the molar ratio of Na₂CO₃ to TiO₂ was 1:6. The mixture was put into a platinum crucible, heated in an electric muffle furnace with a rate of 200 K h⁻¹ to 1273 K, held at this temperature for 10 h, and then cooled at a rate of −100 K h⁻¹ to 773 K, followed by being naturally cooled to room temperature in the furnace. This sample is referred to as NTO(SSR). Several Na₂Ti₆O₁₃ samples were prepared also in a flux method [27]. The precursors, Na₂CO₃ and TiO₂, and a flux reagent such as NaCl, NaF, NaBr, NaI (Nacalai Tesque), or KCl (Kishida), were mixed for 20 minutes in an alumina mortar and heated in the same temperature program as mentioned above, and the solute (Na₂Ti₆O₁₃) to the flux in the molten mixture was 1:1. The obtained powder was thoroughly washed with hot water (353 K) for 15 minutes and filtrated, and this washing step was repeated 3 times to remove the remained residual salt, then dried at 323 K for 24 hours. The Na₂Ti₆O₁₃ samples prepared by different fluxes are referred to as NTO(flux) such as NTO(NaCl).

Ag cocatalyst was loaded on the sodium titanate samples by a photodeposition method (PD), a chemical reduction method (CR), or an impregnation method (IMP). In the PD method, 1 g of the Na₂Ti₃O₇ or Na₂Ti₆O₁₃ sample in 150 mL of ion-exchanged water was irradiated using a 300 W xenon lamp (PE300BUV) for 30 minutes with magnetically stirring. Then 50 mL of CH₃OH and a required amount of AgNO₃ were added and the suspension was stirred for 30 minutes without irradiation. Based on the loading amount of Ag over different samples, it was irradiated for 2-6 hours, where the light intensity was measured to be 22 mW cm⁻² at 254 ± 10 nm in wavelength. The obtained suspension was filtrated and washed by hot water (353K) 3 times, and dried at 323 K for 24 hours. In the CR method, the Na₂Ti₆O₁₃ sample (1g) was suspended in 50 mL of an aqueous solution of AgNO₃ (0.295 M), followed by the dropwise addition of an aqueous NaPH₂O₂ solution (0.295 M, 315 μL) into the suspension. After stirring the mixture at 353 K for 2 h, it was washed by ion-exchanged water, and it was filtered and dried at room temperature. In the IMP method, 1 g of the Na₂Ti₆O₁₃ sample was suspended in 100 mL of ion exchange water with the desired amount of AgNO₃, followed by evaporation at 353 K to remove water, drying for 24 h at 373 K, and calcination at 723 K for 2 h in air. The Ag-loaded NTO(NaCl) samples are referred to as Ag/NTO(NaCl,PD), Ag/NTO(NaCl,CR), and
Ag/NTO(NaCl,IMP).

2.2. Characterization

The loading amount of Ag on the samples was evaluated by X-ray fluorescence analysis with an EDX-8000 (Shimadzu) using an experimentally obtained calibration curve. The crystal structure of the samples was determined by powder X-ray diffraction with a Lab X XRD-6000 (Shimazu). Morphologies of the samples were observed by scanning electron microscopy (SEM) image with a JSM-890 (JEOL). Diffuse reflectance (DR) UV-Vis spectrum was recorded by a V-670 (JASCO). The bandgap was estimated according to the Davis–Mott equation using the Kubelka–Munk function $F(R\infty)$ obtained from the diffuse-reflectance spectrum [42]. The Brunauer–Emmett–Teller (BET) specific surface area was calculated from the amount of N$_2$ adsorption at 77 K, which was measured by a Monosorb (Quantachrome).

2.3. Photocatalytic reaction test

The photocatalytic activity tests of CO$_2$ reduction with water were carried out using a bubbling-flow system with an inner-irradiation-type reaction vessel [24], which were the most different points of the reaction conditions from those in the previous study [27]. The Ag-loaded sodium titanate sample (0.2 g) was dispersed in ion-exchanged water (350 mL) containing 0.5 M NaHCO$_3$ by magnetically stirring. Then, CO$_2$ was bubbled into the solution at a flow rate of 15 mL min$^{-1}$ without irradiation for 1.5 hours. The photocatalytic reaction was conducted using a 100 W high-pressure mercury lamp with 44 mW cm$^{-2}$ measured at 254 ± 10 nm in wavelength. The reaction temperature was 290 K. The evolution rate of the products (CO, H$_2$, and O$_2$) in the outlet from the reactor were determined by using an on-line gas chromatograph (Shimadzu, GC-8A, TCD, Shincarbon ST column, argon carrier).

2.4. Evaluation of oxidative byproducts in aqueous solution

To know whether hydrogen peroxide (H$_2$O$_2$) was formed or not, the aqueous solution after the photocatalytic reaction test was sampled and the conventional starch-iodine test was examined using KI reagent and starch. Assuming that all photoformed holes were consumed to produce H$_2$O$_2$ in this system, the equivalent H$_2$O$_2$ reagent (31.9 μmol L$^{-1}$) was prepared and used as a reference.

To confirm the formation of hypochlorous acid (HClO) in the aqueous solutions after the photocatalytic reactions, a conventional DPD test was employed [39,43,44]. When the $N$-$N$-diethyl-$p$-phenylenediamine (DPD) reagent is added to a solution containing HClO, a red oxidant of DPD rapidly formed through the stoichiometric reaction. The DPD solution was prepared by dissolving 0.1 g of the DPD reagent (Wako), which is a mixture of DPD sulfate and Na$_2$SO$_4$, in 5.0 mL of a phosphate buffer solution (pH = 6.5). 1.0 mL of the prepared DPD solution was added to 5.0 mL of the sample solution, and was shaken for 20 s. The transmittance spectrum was measured immediately using a UV-vis spectrometer (JASCO V-670) in a transmission mode.
3. Results and discussions

3.1. Sodium titanates

Two types of sodium titanates, sodium trititanate (Na$_2$Ti$_3$O$_7$) and sodium hexatitanate (Na$_2$Ti$_6$O$_{13}$), were fabricated. Fig. 1 shows XRD patterns of the obtained samples with those from the database (ICSD#15463 for Na$_2$Ti$_3$O$_7$ [45] and ICSD#23877 for Na$_2$Ti$_6$O$_{13}$ [46]) as references. The patterns of the Na$_2$Ti$_3$O$_7$(SSR) sample and the Na$_2$Ti$_6$O$_{13}$ samples (Fig. 1a, 1b, and 1c) were consistent with the reference data, respectively. Since no diffraction lines corresponding to other impurity phases were observed, it was confirmed that two types of pure sodium titanates, Na$_2$Ti$_3$O$_7$ and Na$_2$Ti$_6$O$_{13}$, were correctly fabricated. It is noted that the distribution of the diffraction line intensity of the NTO(NaCl) sample prepared with the NaCl flux are different from that of the NTO(SSR) sample and the reference. And the intensities of some diffraction lines were also quite different, for example, the intensity of the line at 11.86° corresponding to (200) plane in the pattern of the NTO(NaCl) sample was much higher than that for the others, implying that the Na$_2$Ti$_6$O$_{13}$ crystals prepared by the flux method have an anisotropic morphology.

![XRD patterns of the samples](image)

Fig. 1. XRD patterns of the samples; (a) the Na$_2$Ti$_3$O$_7$(SSR) sample, (b) the NTO(SSR) sample, and (c) the NTO(NaCl) sample. Silicon powder mixed to calibrate the angle showed a diffraction indicated by a closed triangle. The patterns from the ICSD database are also shown, ICSD#15463 for Na$_2$Ti$_3$O$_7$ and ICSD#23877 for Na$_2$Ti$_6$O$_{13}$.

Fig. 2 shows the SEM images. The Na$_2$Ti$_3$O$_7$(SSR) sample consisted of small roundish rod-like or roundish particles with almost uniformed size less than 1 μm (Fig. 2a), while the NTO(SSR) sample showed more irregular shape of particles, most of which were smaller than 1 μm but some of which were much larger (Fig. 2b). On the other hand, the NTO(NaCl) sample consisted of anisotropic rod-like crystal structure covered with facets and crystal size was larger than 2 μm (Fig. 2c). This confirms that the molten salt can enhance the crystallization and regulate the
crystal growth of each facet to form the unique morphology. It is known that this effect depends on the property of the cation such as Na⁺, K⁺ in the molten flux [27].

Fig. 2 SEM images of the prepared samples; (a) the Na₂Ti₃O₇ (SSR) sample, (b) the NTO (SSR) sample, and (c) the NTO (NaCl) sample.

As shown in Table 1, the specific surface areas of the Na₂Ti₃O₇(SSR) sample, the NTO(SSR) sample and the NTO (NaCl) sample were measured to be 2.58, 3.91, and 5.17 m²g⁻¹, respectively. Furthermore, Fig. 3A shows DR UV–visible spectra of these samples, which give information for the electronic band structure. The Na₂Ti₃O₇(SSR) sample showed an absorption band less than 320 nm in wavelength, while the NTO(SSR) and NTO(NaCl) samples showed the absorption edge at 350.7 nm and 351.2 nm, respectively. The estimated bandgap for the Na₂Ti₃O₇(SSR) sample was 3.88 eV, which was slightly bigger than 3.54 eV for the NTO(SSR) sample and 3.53 eV for the NTO(NaCl) sample.

Table 1 Properties of the prepared Na₂Ti₃O₇ and Na₂Ti₆O₁₃ samples.

<table>
<thead>
<tr>
<th>Entry</th>
<th>Sample Preparation method</th>
<th>Crystal phaseᵃ</th>
<th>Specific surface areaᵇ / m² g⁻¹</th>
<th>Bandgapᶜ / eV</th>
<th>Loading amount of Agᵈ (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Na₂Ti₃O₇(SSR) SSR</td>
<td>Na₂Ti₃O₇</td>
<td>2.58</td>
<td>3.88</td>
<td>0.99</td>
</tr>
<tr>
<td>2</td>
<td>NTO(SSR) SSR</td>
<td>Na₂Ti₆O₁₃</td>
<td>3.91</td>
<td>3.54</td>
<td>0.99</td>
</tr>
<tr>
<td>3</td>
<td>NTO(NaCl) Flux method</td>
<td>Na₂Ti₆O₁₃</td>
<td>5.17</td>
<td>3.53</td>
<td>0.93</td>
</tr>
</tbody>
</table>

ᵃ Estimated by X-ray diffraction at 10–80 degree. ᵇ Estimated by BET method. ᶜ Estimated from DR UV-vis spectra. ᵈ Estimated by XRF measurements.

XRF measurements confirmed the loading amount of the Ag cocatalyst on these three samples by the PD method to be almost 1 wt% of Ag cocatalyst on each sample as expected (Table 1). In the DR UV-vis spectra of the Ag-loaded samples (Fig. 3B), the surface plasmon resonance (SPR) band of the Ag nanoparticles was similarly observed for the Ag/NTO(SSR) sample and the Ag/NTO(NaCl) sample, while no such band was observed for the Ag/Na₂Ti₃O₇ sample. This difference would originate from the state of the Ag species added on the sodium titanate samples. For the Ag/Na₂Ti₃O₇ sample, the bandgap was shifted from 3.88 eV to 3.49 eV, where the Ag⁺ cations could be easy inserted and exchange to the Na⁺ between the titanate layered structure of the Na₂Ti₃O₇ sample. This indicated that
Ag nanoparticles exist only on the NTO samples.

Fig. 3 shows the results of the photocatalytic reaction tests, i.e., the time course of the formation rates of the products and the selectivity toward CO evolution \( (S_{CO}) \) in the photocatalytic conversion of CO\(_2\) with water over the three Ag-loaded samples. The three samples exhibited the photocatalytic activity to produce CO, O\(_2\) and H\(_2\). First, when compared the two samples prepared by the SSR method, the Ag/Na\(_2\)Ti\(_3\)O\(_7\)(SSR) and Ag/NTO(SSR) samples (Figs. 4a and 4b), the most obvious difference between them was the product selectivity, i.e., the former exhibited high selectivity to form H\(_2\) and the latter predominantly produced CO with high selectivity such as \( S_{CO} = 75\text{–}90\% \). These samples have some different properties such as compositions, crystal structures, band structure and also the state of the Ag cocatalyst. Especially, the former exhibited no obvious SPR band while the latter clearly showed, suggesting the Ag species exist as a dispersed or oxidized form in the former sample and as nanoparticles in the latter sample. Considering the well-known function of the Ag nanoparticles as a cocatalyst to produce CO, it seems reasonable that the Ag/Na\(_2\)Ti\(_3\)O\(_7\)(SSR) sample without any Ag nanoparticles cannot produce CO and the Ag/NTO(SSR) sample can produce CO selectively.

Next, among the two Na\(_2\)Ti\(_6\)O\(_{13}\) samples, the Ag/NTO(NaCl) sample prepared by a flux method gave the higher formation rate of both CO and H\(_2\) than the Ag/NTO(SSR) sample (Fig. 4c). This shows an advantage of the flux method to give the active Na\(_2\)Ti\(_6\)O\(_{13}\) photocatalyst for this photocatalytic reaction. The CO formation rate over the present Ag/NTO(NaCl) sample with 1 wt% of Ag cocatalyst in the present reaction condition was 3.3 \( \mu \)mol h\(^{-1}\) at 5 h, which is ca. 10 times higher than that over the best Ag/NTO sample in the previous study \([23]\). The both Ag/NTO(NaCl) and Ag/NTO(SSR) samples showed similarly high CO selectivity (Fig. 4b and 4c). This is consistent with the fact that the Ag cocatalyst nanoparticles as the cocatalyst on the NTO surface showed similar SPR band in the UV-vis spectra (Fig. 3).

On both the Ag/NTO samples prepared by the SSR method and the flux method, the formation rate of CO and H\(_2\) decreased and the \( S_{CO} \) increased with irradiation time, which would be due to the aggregation of the deposited Ag particles as mentioned later and also in the previous study \([23]\). The O\(_2\) evolution was also observed in the current reaction condition although the ratio of consumed electron and holes was not consistent, \( R(e^-/h^+)\sim 1 \), which will be
discussed later.

According to the results in this section, further studies were carried out on the Ag/NTO samples prepared by the flux method.

![Graph showing formation rates of CO, H2, and O2, and selectivity toward CO evolution](image)

Fig. 4. Formation rates of CO (red triangles), H2 (black squares), and O2 (blue circles), and the selectivity toward CO evolution (open circles) in the photocatalytic conversion of CO2 with water over the prepared samples loaded with 1 wt% Ag cocatalyst loaded by the PD method, (a) the Ag/Na2Ti3O7(SSR) sample, (b) the Ag/NTO(SSR), and (c) the Ag/NTO(NaCl) sample.

### 3.2. Sodium hexatitanate prepared by the flux method

It was well known that molten salt as a flux can regulate the crystal growth to provide a unique morphology of the crystals. In the previous study, some kinds of the salts such as LiCl, NaCl, KCl, and CaCl2 were examined for the preparation of NTO samples, and it was found that the LiCl and CaCl2 salts are not suitable for the preparation of Na2Ti6O13 phase while only the KCl and NaCl fluxes can promote the formation of the hexagonal rod-like crystals [27]. In order to figure out the effect of NaCl flux, another series of experiments were also conducted in this study. Four catalysts were prepared in the flux method using various fluxes such as NaF, NaCl, NaBr, and NaI. Pure Na2Ti6O13 crystals were confirmed by the XRD patterns (Fig. 5) for these NTO samples except for the NTO(NaI) sample. It is noted that the distribution of the diffraction line intensity of the NTO(NaI) sample are different from those of the NTO(NaCl) and NTO(NaBr) samples and the reference, especially the ratio the intensity of the line at 11.86° corresponding to (200) plane and 14.12º corresponding to (-201) plane. In the UV spectra (Fig. S1), the NTO(NaF) sample exhibited a small shift of the absorption edge, and the NTO(NaBr) sample showed small absorbance of visible light. Furthermore, the SEM images show that the morphologies of NTO with different fluxes were quite different from each other, and it is notable that the NTO(NaCl) shown in Fig. 2c gave much clear morphology with uniform rod-like particles than others shown in Fig. 6. This may be due to the effect of the difference in the electronegativity of the halogen atoms, i.e., 3.98, 3.16, 2.96, and 2.66 [47] for F, Cl, Br and I, respectively, or the melting point of the fluxes, i.e., 1266, 1074, 1028, and 933 K for NaF, NaCl, NaBr, NaI, respectively. Although
the reason could not be clearly mentioned, it is the fact that the NaCl is quite suitable for the fabrication of the rod-like NTO particles.

The results of the photocatalytic activity tests of these samples loaded with Ag cocatalyst were shown in Fig. 7. It is quite clear that the sample prepared with NaCl exhibited the highest activity for CO evolution. It is also noted that the \( R(e^-/h^+) \) with the sample prepared with NaCl was 4.6 at the initial reaction stage, which is much higher than other samples. This suggests that the NaCl flux may be beneficial to the preparation of the photocatalyst for CO production but the chloride anion (Cl\(^-\)) may be not good for sufficient oxygen evolution due to the reactivity to hole as discussed later.

In literature, it was reported that a mixture of two fluxes could enhance the activity of Sr\(_2\)KTa\(_5\)O\(_{15}\) photocatalyst [48], which motivated us to fabricate five samples with the different mixing ratio of NaCl and KCl fluxes in this study (Fig. S2). The results are mentioned in the ESI. The conclusion is, however, that the sample prepared with the pure NaCl flux exhibited the best photocatalytic activity among the examined NTO samples (Fig. S3).

In this section, it was concluded the NTO sample prepared by a flux method with the pure NaCl flux is the optimized one for the reduction of CO\(_2\) in the improved reaction condition. It was clarified that both the use of flux and the property of the flux much affected the shape of the particles and also the photocatalytic activity.

![XRD patterns of the prepared NTO samples with different flux; (a) NaF, (b) NaCl, (c) NaBr, (d) NaI, and ICSD#23877 for Na\(_2\)Ti\(_6\)O\(_{13}\). Silicon powder mixed to calibrate the angle showed a diffraction indicated by a closed triangle.](attachment://Fig. 5.png)
3.3. Ag cocatalysts

The Ag cocatalyst has been employed for the CO production in the heterogeneous photocatalytic reduction of CO2 with water. The loading method and loading amount of the Ag cocatalyst have been reported to affect the state of Ag nanoparticles on the photocatalyst and thus the photocatalytic activity and selectivity [20,49–53]. In the current study, the effects of the loading amount of Ag cocatalyst and loading method on the photocatalytic activity of the NTO(NaCl) samples for CO2 conversion were investigated.

Fig. 8 shows the SEM images of the samples with different loading amount of the Ag cocatalyst prepared by the PD method. It was quite clear that the size of Ag particles became bigger with increasing the loading amount. The particle size of Ag over the 0.1wt% Ag/NTO(NaCl,PD), 0.5wt% Ag/NTO(NaCl,PD) and 1.0wt% Ag/NTO(NaCl,PD)
samples were ca. 12–18 nm, 15–25 nm and 21–32 nm respectively. The Ag particles aggregated when the loading amount was over 1.0 wt%. In the DR UV-vis spectra Fig. S4, with the increase of the Ag loading amount, the band due to the Ag nanoparticles became larger and red-shifted, suggesting that the size of the deposited Ag particles increased, which consistent with the phenomenon observed in the SEM images.

Fig. 8. SEM images of the Ag/NTO(NaCl,PD) samples with different loading amount of Ag, (a) 0.1 wt%, (b) 0.5 wt%, (c) 1.0 wt%, (d) 2.0 wt%, (e) 3.0 wt%, and (f) 4.0 wt%, prepared by the PD method.

Fig. 9 shows the formation rate of the products during the photocatalytic conversion of CO$_2$ with water over the Ag/NTO(NaCl,PD) samples with different loading amount of Ag cocatalyst. Both the CO production rate and the $S_{CO}$ increased and with increasing the loading amounts from 0 wt% to 1.0 wt%, suggesting that the Ag cocatalyst enhances the CO formation and leads to decrease the active sites for H$_2$ evolution on the NTO surface. As a result, 1.0 wt% of Ag cocatalyst was the optimized mounts for providing the highest CO formation rate (4.6 μmol h$^{-1}$) and the highest selectivity toward CO evolution (74 %). On the other hand, the photocatalytic activity decreased above 1.0 wt% while $S_{CO}$ became constant to be 60%. This would be related to the aggregation of the Ag particles observed in the SEM images (Fig. 8), i.e., the surface area of Ag cocatalyst and its contact to the NTO surface would decrease due to the aggregation.
The formation rates of CO (red bar), H₂ (black bar), and O₂ (blue bar) and the S_CO (open circles) in the photocatalytic conversion of CO₂ over the Ag/NTO(NaCl,PD) samples with different loading amount of Ag cocatalyst. The formation rates were recorded after 1 hour’s irradiation. Photocatalyst powder: 0.2 g, reaction solution volume: 350 mL, additive: 0.5 M NaHCO₃, Ag loading method: photodeposition method, CO₂ flow rate: 15 mL min⁻¹, light source: 100-W high-pressure Hg lamp.

The SEM images of the 1 wt% Ag/NTO samples prepared by three methods are shown in Fig. 10, which were taken before and after the photocatalytic reaction tests. As synthesized, it was discovered that both the size of the Ag nanoparticles and their surface distribution on the three samples were quite different from each other (Figs. 10a, 10b, and 10c). When it was loaded by the PD method, some part of Ag nanoparticles with a size of 21–32 nm were mainly stabilized on the long facets of the rod-like crystals (Fig. 10a). During the photodeposition of Ag nanoparticles, Ag⁺ cations are reduced by the photoexcited electron, meaning that the long facets would be the reductive facets that provide electrons predominantly. On the sample deposited by CR method, many fine Ag nanoparticles were more uniformly scattered on all the facets of the NTO crystals with the smallest size (10–15 nm) among the three samples (Fig. 10b). In this method, the adsorbed Ag⁺ cation would be chemically reduced to form Ag nanoparticles on each facet. The IMP method provided a little larger Ag particle with a size of 70–75 nm (Fig. 10c). Probably, the procedure of drying up and calcination would make the Ag⁺ cations and Ag oxide species get together to form larger silver oxide particles. As shown in Fig. 11A, this different phenomenon was also consistent with the DR UV-vis spectra of three samples. Characteristic bands assignable to Ag nanoparticles were observed for the Ag/NTO(NaCl,CR) sample and the Ag/NTO(NaCl,PD) sample, but not for the Ag/NTO(NaCl,IMP) sample. And compared with the Ag/NTO(NaCl,PD) sample, the band over Ag/NTO(NaCl,CR) was a bit blue shifted, which indicated smaller size of Ag particles were fabricated by the CR method as shown in the SEM images (Figs. 10a and 10b).
Fig. 10 SEM images of the Ag/NTO(NaCl) samples prepared by (a, d) PD, (b, e) CR, and (c, f) IMP methods, (a–c) before and (d–f) after the photocatalytic reaction tests for 7 h.

After the photocatalytic reaction tests, however, the size and the distribution of the Ag particles became quite similar to each other (Figs. 10d–f), and almost similar bands assignable to Ag cocatalyst were observed for these samples (Fig. 11B). These results indicate that the Ag particles of these samples aggregated. In details, in the Ag/NTO(NaCl,PD) and Ag/NTO(NaCl,CR) samples, the particle sizes of the Ag nanoparticles became slightly larger, while the silver Ag particles were reduced and dispersed on the Ag/NTO(NaCl,IMP) sample. This means that the rearrangement of the Ag atoms takes place through the Ag⁰/Ag⁺ redox cycles as proposed in the previous study [23], and this would be one of the reasons for the variation of the production rates with time during the photocatalytic reaction tests mentioned below.

Fig. 11. DR UV- spectra of the Ag/NTO(NaCl) samples loaded with 1 wt% Ag by the PD, CR, and IMP methods,
Fig. 12 shows the formation rates of CO, H$_2$, and O$_2$ in the photocatalytic reaction tests for the conversion of CO$_2$ using the Ag/NTO(NaCl) samples, on which 1 wt% of Ag cocatalyst was loaded by the PD, CR, and IMP methods. Among the photocatalysts, the Ag/NTO(NaCl,PD) and Ag/NTO(NaCl,CR) samples exhibited higher photocatalytic activity than the Ag/NTO(NaCl,IMP) sample. And the CO production rate over the Ag/NTO(NaCl,PD) sample was 4.6 μmol h$^{-1}$ after irradiation for one hour, which was higher than the other samples. Since it is proposed that the PD method tends to form the Ag nanoparticles on the reductive facets of the rod-like NTO(NaCl) crystals, the Ag nanoparticles would efficiently receive the photo-excited electrons and promote the photocatalytic reduction of CO$_2$ to produce CO. The CO production rate gradually decreased from the start and became 2.8 μmol h$^{-1}$ after 7 h over the Ag/NTO(NaCl,PD) sample. The aggregation of Ag cocatalyst particles was one responsible reason for this deactivation. As shown in the SEM images (Fig. 10), the Ag particle sizes of the Ag/NTO(NaCl,PD) and Ag/NTO(NaCl,CR) samples became to 25-30 nm after 7 hours’ reaction test, which could decrease the active sites of the Ag cocatalyst surface and the contact to the NTO surface, and thus reduce the photocatalytic activity.

In the previous reaction conditions [27], the formation of O$_2$ was not observed initially. Even though it became actually observable 24 hours later, it was still not enough production rate compared to the formation rates of reductive products. In the present study, the O$_2$ evolution was observed even at the initial stage. However, the stoichiometric formation rate of O$_2$ was still not obtained during the initial period over the Ag/NTO(NaCl,PD) sample, where $R(e^-/h^+)$ was 3.0 after 7 hours’ irradiation. This matter will be discussed in the next section.
3.4. Oxidation reaction

The oxygen evolution reaction (OER) is one of the major challenges in the water splitting as well as CO₂ reduction with water. Many papers have been reported to improve the OER by combined the co-catalysts with some metal oxides such as manganese oxide, and so on [54–59]. And in literature, it is sometimes found that photocatalysts for water splitting produce only H₂ without enough evolution of O₂ (H₂/O₂>2) [31,60]. This phenomenon may be explained by the difficulty of O₂ evolution due to the high overpotential of O₂ formation, which originates from trapping and stabilization of positive charge by the surface sites [28–30]. However, if the consumption of holes does not take place, the reductive reaction to produce CO and H₂ will also stop soon. Since the formation of reductive products underwent over 8 h in Fig. 9, the oxidation reaction should take place. Here, three possibilities are discussed as the reasons for this phenomenon.

The first possibility is that water might be oxidized to hydrogen peroxide [61] (Eq. 8).

\[
2\text{H}_2\text{O} + 2\text{h}^+ \rightarrow \text{H}_2\text{O}_2 + 2\text{H}^+ \quad (8)
\]

To confirm this possibility, the starch iodine reaction test was carried out. It showed a negative result for the solution after the photocatalytic reaction test as described in the supplementary information, i.e., the color change was not observed as shown in Fig. S5. Thus, it can be mentioned that this was at least not the dominant reason.

The second possibility is that residual chloride anion (Cl⁻) on the surface of the NTO(NaCl,PD) photocatalyst, originating from the NaCl flux, might react with the photoformed hole to form hypochlorite (ClO⁻) shown in eq. 9, which competitively consume hole and reduce the oxidation of water to form O₂.

\[
\text{Cl}^- + \text{H}_2\text{O} + 2\text{h}^+ \rightarrow \text{ClO}^- + 2\text{H}^+ \quad (9)
\]

\[
2\text{ClO}^- + \text{hv} (\lambda < 400 \text{ nm}) \rightarrow \text{O}_2 + 2\text{Cl}^- \quad (10)
\]

The DPD tests [39] were conducted for the filtrated solutions after the photocatalytic reaction tests with the Ag/NTO(SSR,PD) sample and the Ag/NTO(NaCl,PD) sample as well as a distilled water as a reference, and the positive results were obtained on the solution separated from the solution with the Ag/NTO(NaCl,PD) sample (Fig. S6). Thus, it was clarified that Cl⁻ residue was oxidized to ClO⁻, which competitively consumed holes to reduce the formation of O₂. The detected amount of ClO⁻, calculated by the standard curve reported in the literature [42], was quite small (0.8 μmol) after 7 hours’ irradiation. This may be due to the decomposition of hypochlorite (ClO⁻) shown in the eq. 10 since the high-pressure mercury lamp was used in the current system. With the increasing concentration of ClO⁻ in the solution, the reaction rate of eq.9 would decrease while the rate of eq.10 would increase. And when the reaction rate of the eq. 10 increased to a half of that of the eq. 9, the stoichiometric amount of O₂ was supposed to be produced in the current system.
Fig. 13. Formation rates of CO (red triangles), H₂ (black squares), and O₂ (blue circles) and the $R(e^-/h^+)$ (open circle in the photocatalytic conversion of CO₂ with water by using (a) the Ag/NTO(SSR,PD) sample and (b) the Ag/NTO(NaCl,PD) sample. Photocatalyst powder: 0.2 g, reaction solution volume: 350 mL, additive: 0.5 M NaHCO₃, Ag loading method: photodeposition method, CO₂ flow rate: 15 mL min⁻¹, light source: 100-W high-pressure Hg lamp.

The third possibility is that the produced O₂ might be adsorbed on the surface of catalysts to some extent especially under photoirradiation, since the photoadsorption of oxygen on TiO₂ was reported [32,33]. Fig. 13 shows the formation rates of CO, H₂, and O₂ and the consumed $e^-/h^+$ ratio, $R(e^-/h^+)$, in the photocatalytic conversion of CO₂ over the two samples, Ag/NTO(SSR,PD) and Ag/NTO(NaCl,PD). Even in the case of the reaction test with the Ag/NTO(SSR,PD) sample, the O₂ evolution increased with irradiation time, and the $R(e^-/h^+)$ became to be 1.0 after 22.5 hours. Since this sample should be free from chloride anion, the initially insufficient O₂ evolution could be explained by not the oxidation of chloride anion but the photoadsorption of O₂ on the surface of catalysts. After 22.5 hours, the O₂ photoadsorption on the surface would be saturated and no more photoadsorption would take place.

Even for the Ag/NTO(NaCl,PD) samples, the ratio of $e^-/h^+$ became gradually close to be 1.0 after about 50 hours. In this state, not only the O₂ photoadsorption but also the composition of the chloride (Cl⁻) should be taken into consideration. The concentration of hypochlorite (ClO⁻) anion would also become constant in the reaction solution to give O₂ evolution.

As a summary of this section, two possible reasons for the non-stoichiometric amount of O₂ evolution can be proposed as follows, i.e., the oxidation of chloride residues and the photoadsorption of O₂ on the surface.

4. Conclusion

In this study, two remained problems in the previous work, one of which is low activity toward CO evolution and low selectivity, and the other is nonstoichiometric O₂ evolution, were much improved and explained, respectively.

First, it was found that the Ag/NTO sample, with 1.0 wt% of Ag cocatalyst loaded by a photodeposition method, exhibited the highest CO production rate (4.6 μmol h⁻¹) and the highest CO selectivity (74%) among the examined samples in the current reaction conditions. These values are more than 29 times higher production rate and more than 2.7 times higher selectivity than those obtained in the previous study, which would originate from the optimization of the photocatalyst and the improved reaction conditions. In addition, among three Ag/NTO samples, on which 1.0
wt% of the Ag cocatalyst nanoparticles were loaded by different methods, the Ag/NTO sample provided by the PD method gave the highest activity and selectivity for CO₂ reduction. Since the PD method tends to form the metallic Ag nanoparticles on the reductive facets of the rod-like NTO crystals, which can efficiently promote the CO evolution.

Second, although the evolution rate of O₂ was not enough in the induction period, it increased with time and reached to the stoichiometric formation ratio of the oxidative and reductive products after a long time, such as 24 or 50 h, depending on the samples. Furthermore, it was confirmed that the chloride residues and the photoadsorption of O₂ on the surface are responsible for the insufficient O₂ evolution less than stoichiometric ratio among the products at the initial period.

The present comprehensive study has provided some solutions to improve both the CO evolution and the S_{CO} over the Na₂Ti₆O₁₃ sample. Further studies are very necessary both to improve the photocatalytic activity and enhance the O₂ evolution at the initial time.

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