Effects of sol–gel procedures on the photocatalysis of Cu/TiO$_2$ in CO$_2$ photoreduction

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Received 28 January 2003; revised 27 August 2003; accepted 3 September 2003

Abstract

Copper-loaded titania (Cu/TiO$_2$) was synthesized via an improved modified sol–gel process. Photocatalysts were applied to the CO$_2$ photocatalytic reduction and the yield of the major product, methanol, was used to evaluate the photocatalytic performance. Copper precursors and the adding time with sol as well as posttreatments were studied to explore the relationships between the characteristics and the activity of the photocatalysts. The results revealed that Cu/TiO$_2$ prepared from copper chloride and added in the early sol–gel stage was more photoactive than that from copper acetate. Additional H$_2$ reduction of calcined catalysts before the photoreduction CO$_2$ decreased the yield of methanol due to the change of copper dispersion and oxidation state. TPR, XPS, and XAS measurements verified the oxidation state of Cu on Cu/TiO$_2$ catalysts. The results indicated that the primary Cu(I) served as an active site. The zeta potentials of catalysts were measured and compared, showing that a higher positive zeta potential at pH 7 would lead to higher activity. Under 30-h UVC (254 nm) irradiation, the best catalyst gave a methanol yield above 600 µmol/g$_{\text{cat}}$. Switching to UV A (365 nm) resulted in a significant decrease of methanol yield in the range of 10 µmol/g$_{\text{cat}}$.

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Keywords: Cu/TiO$_2$; CO$_2$; Photoreduction; XANES; EXAFS; Sol–gel

1. Introduction

The sol–gel process is suitable for producing composite materials of high purity without multiple steps. In previous work [1], homogeneous, nano-sized, copper-loaded anatase titania was synthesized by the improved sol–gel method. These titania photocatalysts were applied to the photoreduction of carbon dioxide in order to evaluate their photocatalytic performance. Methanol was found to be the primary hydrocarbon product [2].

Various methods exist for solving the problem of the greenhouse gas, carbon dioxide, on earth. One permanent solution is to transfer it chemically into a useful and nontoxic material. Several studies [3–5] have found that CO$_2$ can be transformed into methanol in a photocatalytic reaction. Methanol is an important chemical and convenient liquid fuel.

One crucial constraint on solving the problem of CO$_2$ is that any energy source used should not produce more CO$_2$. Solar energy is one of the best choices because it is clean and inexhaustible. However, the quantum efficiency is so far very low. Huynh et al. [6] reported a solar power conversion efficiency of only 10% in conventional inorganic solar cells; the most advanced but expensive cell has an efficiency of 30%. Moreover, 95% of solar light reaching ground is visible light [7,8]. Consequently, highly efficient photoreduction of CO$_2$ is preferentially desired.

Previous work [2] revealed that the optimum copper loading was 2 wt% due to the highest Cu dispersion. The best operation conditions for CO$_2$ photoreduction, including CO$_2$ pressure, weight of catalysts, and addition of NaOH, have been studied. This work aims to extend the fundamental understanding of the relationships among the sol–gel procedure, the properties of catalysts, and photocatalytic activity. The preparation parameters, such as the copper precursors, adding times, and posttreatments, are under investigation. This understanding can eventually be applied to develop visible-light-responsive photocatalysts, and to control the products selectivity of photoreaction using solar energy.
2. Experimental

2.1. Preparation of Cu/TiO$_2$ catalysts

An improved sol–gel process was used to synthesize copper-loaded titania photocatalysts. Fig. 1 depicts the procedure. Titanium butaoxide, $n$-butanol, and acetic acid were vigorously stirred in a beaker in a temperature and humidity-controlled glove box. Copper precursor was added during the above hydrolysis and polycondensation period. The resulting transparent green sol was dried at 150 °C and calcined at 500 °C in flowing air. The catalyst was then pulverized. Different adding periods, ranging from 0 to 8 h, and copper precursors, either copper chloride or copper acetate, were used. The procedure is detailed in a previous work [1]. Additionally, posttreatments of the resultant catalysts before reaction were also investigated. The H$_2$ reduction conditions were carried out at 300 °C for 3 h in a 5% H$_2$/Ar flow. For comparison, two commercial titania, Merck and Degussa P25, were deposited or impregnated by either the photodeposition (P) or the incipient wetness (I) method to make Cu/P25 and Cu/Merck catalysts, respectively. Table 1 summarizes the preparation conditions and the abbreviated forms of the prepared catalysts.

### Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Procedure</th>
<th>Cu (wt%)</th>
<th>Copper precursor</th>
<th>Adding time (h)</th>
<th>Posttreatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuCl$_2$-x h</td>
<td>Sol–gel</td>
<td>2</td>
<td>CuCl$_2$</td>
<td>$x=0$–8</td>
<td>None</td>
</tr>
<tr>
<td>Cu(NO$_3$)$_2$-y h</td>
<td>Sol–gel</td>
<td>2</td>
<td>Cu(NO$_3$)$_2$</td>
<td>$y=0$–8</td>
<td>None</td>
</tr>
<tr>
<td>CuCl$_2$-x h</td>
<td>Sol–gel</td>
<td>2</td>
<td>CuCl$_2$</td>
<td>$x=0$–8</td>
<td>H$_2$ reduction</td>
</tr>
<tr>
<td>Cu(NO$_3$)$_2$-y h</td>
<td>Sol–gel</td>
<td>2</td>
<td>Cu(NO$_3$)$_2$</td>
<td>$y=0$–8</td>
<td>H$_2$ reduction</td>
</tr>
<tr>
<td>Cu/Merck</td>
<td>P$^a$</td>
<td>&lt; 2</td>
<td>CuCl$_2$</td>
<td>–</td>
<td>None</td>
</tr>
<tr>
<td>Cu/P25</td>
<td>I$^b$</td>
<td>2</td>
<td>CuCl$_2$</td>
<td>–</td>
<td>None</td>
</tr>
</tbody>
</table>

$^a$ P: photodeposition method.

$^b$ I: incipient wetness method.

2.2. Characterization

The specific surface area of the catalyst was measured by N$_2$ adsorption using a Micrometrics ASAP 2010. A diffuse reflectance UV–vis spectrophotometer (Hitachi, U3410) was employed to obtain the UV–vis spectrum. The crystalline phase was identified by X-ray diffractometry (XRD) on a MAC (M03XHF, Material Analysis and Characterization, Japan). X-ray photoelectron spectroscopy (XPS) was performed using a VG Microtech MT500 with an Mg-K$_\alpha$ X-ray source. All binding energies were referenced to oxygen (1s) at 530.7 eV or carbon (1s) at 285.6 eV. A scanning electron microscope (SEM), equipped with an energy-dispersive spectrometer (EDS) LEO 1530 FEG–SEM–EDS, was used to observe the morphology of the catalyst and to measure its elemental ratio. The element distribution was analyzed by measuring the X-rays emitted after an exciting electron beam (15 keV) scanned a small area of a compressed catalyst pellet. In the temperature-programmed reduction (TPR) experiments, a quartz sample holder was loaded with approximately 0.1 g of fresh catalysts. The sample was heated from room temperature to 500 °C in a 5% H$_2$/Ar stream. A flow rate of 30 ml/min and a heating rate of 10 °C/min were used in all TPR experiments.

The X-ray absorption spectra (XAS) of the Cu and Ti K edge for all catalysts were measured at the Wiggler 17C station of the Taiwan Synchrotron Radiation Center in Hsinchu Science-based Industrial Park. A fluorescence mode was used to make the XAS measurement of a small amount of titania-supported copper catalyst. The powder sample was pressed in a sample holder positioned at 45° to the incident X-ray beam in a sample box. The fluorescent X-rays from the sample passed through a Ni filter and reached the detector. The filter was used to avoid the interference from other atoms or incident X-rays. The measurement of titanium was in transition mode directly. The X-ray photon energy varied across and beyond the absorption edge of the measured atom. For Cu, it was in the range from 200 eV below the copper absorption edge at 8979 to 800 eV above it. The intensity of both fluorescent ($I_f$) and incident ($I_o$) X-rays was measured to calculate the absorption coefficient ($\mu$) for the
atoms of interest using the equation, \( \mu_x = I_o/I_f \), where \( x \) is the thickness of the sample. Spectral analysis followed the standard steps of background correction and normalization, before the near-edge absorption structure was determined. Moreover, the spectra of pure \( \text{Cu}_2\text{O} \), \( \text{CuO} \) powder, and \( \text{Cu} \) foil were measured as standard references.

Generally, XAS can be divided into two types of structures superimposed on an X-ray absorption edge. The first structure is the X-ray absorption near-edge structure (XANES), which pertains to energies up to around 50 eV above the absorption edge. The second one extends from around 50 eV above the absorption threshold to several hundred eV above it, and is known as the extended X-ray absorption fine structure (EXAFS). These two kinds of spectra in this work were derived from the WinXAS 2.33 software [9].

Catalyst particles were first spread in distilled water and pumped into a capillary cell to measure zeta potential. Laser beams illuminated the particles in the capillary cell to generate their electrophoretic mobility. The Zeta Sizer, Malvern 2000HS A, recorded the electrophoretic mobility and then related to the Henry equation resulting in the zeta potential as a function of pH value. A pH value ranging from 5 to 12 was applied by adjusting the amount of NaOH added; each pH measurement was made five times.

2.3. Photoreduction of CO\(_2\)

The experiment was carried out in a cylindrical quartz reactor with a capacity of 300 ml. A weight 0.3 g of catalyst powder was suspended in 0.2 N NaOH solution. This condition was optimal according to a previous study [2]. Ultrapure CO\(_2\) from Air Products and Chemicals was bubbled through the reactor for at least 4 h to ensure that all dissolved oxygen was eliminated; then, the irradiation lamp was turned on to start the photoreaction. The illumination system included a mercury lamp (Ultra-Violet Products Inc., USA; 11SC-1), either UVC (254 nm) or UV A (365 nm), in the center of the reactor. The whole system was tightly closed during the irradiation. The temperature, pH value, and dissolved oxygen concentration were monitored continuously. A needle-type probe was inserted into the reactor to withdraw a small liquid sample. After catalyst particles were filtered, the sample was analyzed using a GC-equipped flame ion detector and a 2-m-long Porapak Q column. Analytical results indicated that methanol was the major hydrocarbon.

3. Results

3.1. Characteristics of catalysts

From the previous study [2], 2 wt% \( \text{Cu/\text{TiO}_2} \) catalyst yielded the highest methanol under a 6-h UVC irradiation. In this study, all the copper loadings of the catalysts were maintained at about 2 wt% to investigate other effects. The XRD spectra indicate only the anatase phase of titania, and no significant peaks related to copper phases for any sol–gel-derived \( \text{Cu/\text{TiO}_2} \). The sizes of crystals estimated from the Scherrer equation were all about 20 nm. Different sol–gel procedures did not result in notably different XRD or BET results. The BET surface areas of all the catalysts ranged from 20 to 30 m\(^2\)/g and were less than that of pure TiO\(_2\), 63 m\(^2\)/g. The “CuAc\(_2\)-8 h” catalyst was an exception, with a surface area of 67 m\(^2\)/g, similar to that of pure TiO\(_2\). The micropores of the catalysts were identified from the N\(_2\) adsorption isotherms. Fig. 2 displays the UV–vis spectra that show the influence of copper on the UV–vis absorption. For pure titania (Merck), the absorption is associated with the excitation of the O 2p electron to the Ti 3d level [10]. The absorption edge extends to longer wavelengths for \( \text{Cu/\text{TiO}_2} \), revealing good contact between \( \text{TiO}_2 \) and \( \text{Cu} \) grains [11].

Fig. 3 shows the results of XPS spectra of 2 wt% \( \text{Cu/\text{TiO}_2} \) prepared from various procedures. Because of small Cu loading and high photoionization cross section [12], reference spectra of copper particles, \( \text{Cu}_2\text{O} \) and \( \text{CuO} \), are also shown in Fig. 3 to identify the copper state on the surface of TiO\(_2\). The Cu (2p)-binding energies of \( \text{Cu}_2\text{O} \) were found to be 932.8 and 952.8 eV, respectively. However, the binding energies for CuO were 1 eV above those for \( \text{Cu}_2\text{O} \), at 933.8 and 953.8 eV, respectively. According to the position and the shape of the peaks, regardless of various procedures, the copper on the surface of TiO\(_2\) may exist in multiple-oxidation states but Cu(I) is the primary species even after H\(_2\) reduction.

Table 2 presents the quantitative analysis calculated from the XPS. For 2 wt% Cu loading, the bulk Cu/Ti molar ratio was calculated to be about 0.026, but the quantitative analysis revealed much higher Cu/Ti ratios, estimated from the XPS. XPS can only detect the outer most surface (~ 10 nm)
of a sample. The results indicate that copper was dispersed mostly on the surface of the prepared catalysts. The substitution of copper chloride to copper acetate reduced the Cu/Ti ratio. The catalyst CuCl₂-1 h had the lowest Cu ratio on the surface. For comparison, the Cu/Merck catalyst prepared by the photodeposition method had all copper particles on its surface.

The quantitative analysis of the elements in catalysts was also estimated by EDS analysis. Table 2 also presents relative bulk Cu/Ti molar ratio from EDS. The electron energy (15 keV) of the EDS can detect elements near 1 µm in depth from a sample surface. The results indicate that these values are slightly larger than the calculated bulk value of 0.026. When CuCl₂ was used as a copper precursor, detectable Cl remained in the catalysts. Chlorine resided in the catalysts even after calcination or H₂ reduction. The molar ratio of Cl was calculated based on the area of peaks from EDS. The amount of residual Cl was found to be affected by the adding time of CuCl₂ during the hydrolysis period. The Cl molar ratio reached a maximum ∼0.6% when CuCl₂ was added between 1 and 3 h of hydrolysis. Posttreatment by H₂ reduction at 300 °C for 3 h only eliminated small amount of Cl.

Fig. 4 plots the TPR results of 2 wt% Cu/TiO₂. Pure titania did not consume H₂ in this temperature range. The hydrogen reduction temperatures (T_R) of prepared catalysts via various sol–gel procedures were all near 200 °C. Precursor CuAc₂ had T_R 20 °C lower than that of CuCl₂. However, when copper chloride was added at either 1 or 3 h, a higher T_R was required to reduce these two catalysts. Furthermore, the area under their peaks was smaller than the areas under those of other sol–gel-derived Cu/TiO₂. The TPR of the catalyst used for comparison, Cu/P25 prepared via the incipient wetness method, showed a reduction peak at 280 °C but the area under its peak was larger than the areas under those of all sol–gel-derived Cu/TiO₂.

The XANES of CuO, Cu₂O, and Cu standards were measured for comparison with other samples. A preedge was clearly observed for Cu₂O and Cu standards. For CuO standard, the absorption edge was slightly shifted to higher photon energy. Cu standard gave a split near-edge spectrum. In Fig. 5, results for catalysts with different amounts of copper loading were compared to the Cu XANES spectra. The major Cu(I) oxidation state was concluded according to the absorption edge. A higher Cu loading corresponded to a clearer preedge, especially at a loading above 2 wt%. Catalysts with higher Cu loading appeared to exhibit the characteristic split copper peak. Higher loading resulted in the aggregation of
copper particles and decreased copper dispersion. A copper X-ray diffraction peak of 6 wt% Cu/TiO$_2$ was also observed in a previous study due to aggregation under high Cu loading [2].

Fig. 6 displays the XANES spectra of catalysts prepared with different CuCl$_2$ adding time and the effect of H$_2$ reduction. The position and shape of the absorption edge contributed to the Cu(I) absorption. In addition, Fig. 6 shows some distinguished results. The absorption spectra of CuCl$_2$-1 h and CuCl$_2$-3 h dramatically changed after H$_2$ reduction. The major oxidation state of Cu changed to Cu(0). However, whether or not H$_2$ reduction had occurred, the structures of CuCl$_2$-0 h and CuCl$_2$-8 h were similar.

The FT-EXAFS spectra of CuO, Cu$_2$O, and Cu standards were compared with Figs. 7–9. Fig. 7 displays the effects of various periods of adding CuCl$_2$ or CuAc$_2$ on the FT-EXAFS spectra of Cu/TiO$_2$. Cu(I) particles exhibit two main geometries. One is isolated Cu(I), in which the Cu–O peak is located at around 1.5 Å, and the other is aggregated Cu(I), which contributes to a Cu–O–Cu peak at around 2.6 Å [13,14]. Most of the copper on these catalysts was well-dispersed Cu(I) small particles, resulting in
Fig. 9. Effect of preparation methods on FT-EXAFS spectra. (I, incipient wetness; P, photoreduction; S, sol–gel).

a major peak at 1.5 Å. Except for CuCl₂-3 h catalyst, a very few aggregated Cu₂O (at 2.6 Å) or CuO (at 2.5 Å) particles formed on CuCl₂-0 h, CuCl₂-1 h, and CuCl₂-8 h. Fig. 8 is the FT-EXAFS spectra showing the effect of H₂ reduction. The H₂ reduction only slightly influenced CuCl₂-0 h and CuCl₂-8 h catalysts, the Cu structures of which were similar to that of the well-dispersed isolated Cu(I) particle, even after H₂ reduction treatment. However, lots of Cu(I) particles were reduced to Cu(0) on CuCl₂-1 h and CuCl₂-3 h catalysts, resulting in a characteristic peak at 2.2 Å [14,15].

Another copper precursor of Cu(CH₃COO)₂ was used to prepare 2 wt% Cu/TiO₂ catalysts. Fig. 8 shows the major characteristic peak of r-CuAc-0 h at 1.5 Å, to which the isolated Cu(I) particles contributed. However, the reduced r-Cu(Ac)₂-8 h gave most Cu(0) species resulting in a major peak at 2.2 Å.

Fig. 9 compares the FT-EXAFS spectra for three preparation methods using CuCl₂ precursor. The spectra indicate that 2 wt% Cu/TiO₂ by these methods all resulted in isolated Cu(I) particles. However the sol–gel method still gave less Cu aggregation (a smaller peak at 2.6 Å) than other methods.

The surface potential of a catalyst is a major factor that controls the stability of its aqueous suspension. The point of zero charge (pzc) is a critical pH value in an aqueous solution. At this pH value, the charge on particles is zero and the particles favorably aggregate. Particles can be well dispersed at either higher surface potential or the working pH value far from the pzc. The catalysts particles were suspended in an aqueous solution during the photocatalytic reaction. The influence of pH value on zeta potential was studied for various catalysts. The isoelectric point of most copper-loaded titania in distilled water is below pH 7, similar to that of pure TiO₂ and Degussa P25. Within this pH range, all catalysts, except CuCl₂-0 h and CuAc₂-8 h, carried negative zeta potential, indicating that their surface was negative. The zeta potentials of CuCl₂-0 h and CuAc₂-8 h catalysts were positive at pH 7.

3.2. Photocatalytic activity

Fig. 10 presents the CO₂ photoreduction results for various catalysts. Methanol yield was used to evaluate the performance of the catalysts since it was the major hydrocarbon product. After 30 h of UVC (254 nm) irradiation, the best four catalysts were, in order, CuCl₂-0 h, CuAc₂-8 h, CuCl₂-8 h, and CuAc₂-0 h, with a maximum methanol yield above 600 µmol/gcat. The H₂-reduced catalysts would decrease the methanol yields. An increase wavelength of UV significantly decreased the methanol yield. The methanol yields were only in the range of 10 µmol/gcat under 30 h of irradiation with a UV A (365 nm) lamp, as shown in Fig. 10. The production of methanol by commercial TiO₂ (Degussa P25) under UVA irradiation was undetectable.

3.3. Sol–gel process

Fig. 11 plots the time dependence of pH and temperature during the sol–gel process, indicating that the variation of temperature changes. In the sol–gel process used herein, titanium butoxide was mixed in sequence with butanol and acetic acid. Initially, the temperature of titanium butoxide was about 28 °C; then the temperature quickly increased to 35°C as two solutions were mixed in. Thereafter, the temperature fell to 31 °C within 1 h, increasing again to nearly
33 °C in the third hour. Finally the temperature monotonically decreased and remained stable at near 31.5 °C to the end of the hydrolysis.

4. Discussion

4.1. Copper state and dispersion

The catalytic activity of Cu/TiO\textsubscript{2} in CO\textsubscript{2} photoreduction was related to the preparation and properties of the catalysts. The XPS results in Fig. 3 show that the shape and the position of the Cu 2p-binding energy indicated a combination of Cu(II), Cu(I), and Cu(0) states. Cu(I) was the primary chemical species. Moreover, the difference between the quantitative analysis results obtained by XPS and those obtained by EDS, shown in Table 2, indicates that Cu was near the surface because of the much higher Cu/Ti ratio in the XPS results.

A comparison of Cu absorption edge and near-edge structure between standards and catalysts gave additional evidence of the major Cu(I) state on the catalysts. Furthermore, the extended XAS data analysis elucidated the environments of the Cu atoms. The major peaks in the EXAFS spectra, at 1.5 and 2.2 Å, are characteristic of the isolated Cu(I) and Cu(0) particles, respectively. Aggregated Cu(I) also contributes partially to a peak at 2.6 Å. The derived atomic contribution to the X-ray absorption peak at 2.6 Å indicates notable aggregation of copper particles in the catalysts. The catalyst using copper acetate led to a higher Cu dispersion than CuCl\textsubscript{2}, which exhibited a lower Cu/Ti ratio in the XPS results.

4.2. Effects of sol–gel procedure on copper state

Either CuAc\textsubscript{2} or CuCl\textsubscript{2} precursor added at the beginning with titania sol yielded similar resultant Cu states, that is, of well-dispersed small Cu(I) particles. Adding CuCl\textsubscript{2} at the end of hydrolysis yielded no significantly different results from those adding it at the beginning, while significant differences arose adding it within 1 to 3 h of hydrolysis. When CuCl\textsubscript{2} was added at 1 or 3 h, a high TR was required to reduce both catalysts. Furthermore, the areas of their peaks were smaller than those for other sol–gel-derived Cu/TiO\textsubscript{2}.
Some copper was presumed to be covered by the TiO$_2$ surface [17].

Fig. 11 plots the temperature and pH during the sol–gel process. The first high-temperature peak was associated with the exothermic esterification of butanol and acetic acid. The water released by esterification was then progressively consumed by in the hydrolysis of titanium butaoxide, which is an endothermic reaction, so the temperature decreased gradually. Extra butanol was released during the hydrolysis so that further esterification proceeded, consuming acetic acid. Consequently, the temperature increased again and the pH value increased until the third hour. The composition of the solution at different times changed the solubility and distribution of the copper precursor. CuCl$_2$ added between 1 and 3 h could not be uniformly dispersed so the Cu dispersion of the resultant Cu/TiO$_2$ on the surface was poorer and the interaction with TiO$_2$ was weaker. This fact also explains why the H$_2$ could further reduce Cu(I) to Cu(0), resulting in the aggregation of Cu. Furthermore, the lower Cu dispersion and the weaker interaction caused more residual Cl to remain in the catalysts and increased the H$_2$ reduction temperature ($T_h$), as shown in Fig. 4.

4.3. Major factors controlling CO$_2$ photoreduction

Under the conditions for CO$_2$ photoreduction applied in this work, adding NaOH at the beginning of the reaction resulted in a pH value of 12. After CO$_2$ was bubbled into the solution, saturated solubility was reached and the pH value dropped to 7. All catalysts in the reaction solution were far away from the pzc and were well dispersed. Comparing the variation of zeta potential within this range, most catalysts carried a negative zeta potential, indicating that the surfaces of the catalysts, except CuCl$_2$-0 h and CuAc$_2$-8 h, were negative. CuCl$_2$-0 h and CuAc$_2$-8 h had a positive zeta potential at pH 7.

In Fig. 10, the catalysts that performed well in CO$_2$ photoreduction under UVC irradiation were in the order CuCl$_2$-0 h, CuAc$_2$-8 h, and CuCl$_2$-8 h, which was also consistent with the positive zeta potential at pH 7. The positive zeta potential indicated positive surface charge on the surface of a catalyst. The positive surface enhanced the adsorption of carbonate ions and strengthened the interaction between TiO$_2$ and Cu, improving the photoactivity [7,17]. The surface area of CuAc$_2$-8 h was higher than that of any other catalyst (Table 2) and was believed to facilitate its performance. The photoactivity decreased somewhat after H$_2$ reduction. Based on the FT-EXAFS results, the relative amount of Cu(I) decreased after H$_2$ reduction. Therefore, the active sites for CO$_2$ photoreduction were isolated Cu(I) species. However, the Cu(0) state might also induce the production of other hydrocarbons, as revealed by some experimental results.

5. Conclusions

In this work, the specific surface area and the UV–vis spectra were similar for all investigated catalysts. In our improved sol–gel process and under calcination conditions, small copper particles can be well dispersed on the surface of anatase titania. Copper loaded on the surface enhanced the photoactivity. According to XAS and XPS analysis, the oxidation state of Cu(I) was suggested to be the active species for CO$_2$ photoreduction. Higher copper dispersion and smaller copper particles on the titania surface correspond to a greater improvement in the performance of CO$_2$ photoreduction. The greater the aggregation of copper particles, the higher the H$_2$-reducing temperature required and the larger amount of chlorine resided on catalysts. The choice of a CuCl$_2$ precursor increased the Cu dispersion over that obtained with a copper acetate precursor. The post H$_2$ reduction only slightly affected the Cu oxidation state and caused little aggregation of any sol–gel-derived catalyst, except in the case of any catalyst for which a Cu precursor was added at an inappropriate time during hydrolysis. A positive zeta potential at pH 7 promotes the photoactivity of CO$_2$ photoreduction.

Methanol was favorably produced on Cu/TiO$_2$ catalysts in an aqueous solution of CO$_2$ under UV irradiation. The transformation of photo to chemical energy by a catalyst provides a more efficient way than performed by green plants. The results demonstrate that the improved sol–gel method is a promising technique for preparing such photocatalysts.

Acknowledgments

The authors thank the National Science Council of Taiwan, the Republic of China, for financially supporting this research under Contract NSC-90-2214-E-002-038. The authors also thank Dr. Jyh-Fu Lee of the Wiggler 17C station of the Taiwan Synchrotron Radiation Center and Ms. Chaoling Lai of the Surface Analysis Lab at the National Taiwan University for their effort in instrumental analysis.

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