Effect of Copper Species in a Photocatalytic Synthesis of Methanol from Carbon Dioxide over Copper-doped Titania Catalysts

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Abstract: Synthesis of methanol from CO₂ by a photocatalytic reduction process over copper-doped TiO₂ has been investigated. It has been reported in the literature that copper species could enhance methanol production in a photocatalytic reduction of CO₂. However, which species of copper (e.g. Cu^{0}, Cu^{I}, or Cu^{II}) and how do the copper species play a role still unclear. In this paper we will report and discuss our finding that each copper species play a different role. By using modified-impregnation method in preparing copper/TiO₂, it was revealed that CuO is the copper species, which resulting much better enhancement in methanol production than the others. The activation energy (E_a) for TiO₂ Degussa-P25 and 3%CuO/TiO₂ is ca. 26 and 12 kJ/mol, respectively. The lower E_a of 3%CuO/TiO₂ may suggest a catalytic role of copper species that increase the methanol production. Furthermore, an in situ FTIR study revealed appearance of formate and methoxy intermediates on the CuO/TiO₂ surface, while none if the undoped TiO₂ catalyst was employed. This may indicate a catalytic nature of copper species as an active site to give methanol. Enhancement of product yield was also observed indicates that the copper species act as an electron trapper and prohibit electron-hole recombination.

Key words: Photocatalytic reduction %Titanium dioxide %Copper species %CO₂ %Methanol

INTRODUCTION

The continuously increasing CO₂ level into the atmosphere is one of the most serious problems with regard to the greenhouse effect. Photocatalytic reduction of CO₂ become in the future an alternative solution not only for environmental problems caused by CO₂ emission, but also for finding ways to maintain hydrocarbon resources which now on are being dominated by fuel and natural gas. Methanol as one of the main product of CO₂ photoreduction can be transformed into other useful chemicals and used as fuel-like renewable energy [1]. Much research has been reported that CO₂, in the present of water, can be photocatalytically converted to methane, methanol, or other hydrocarbons over TiO₂ or copper doped TiO₂ [1-5], CoPc loaded TiO₂ [6] and mixed oxide-based photocatalyst [7,8]. It is likely that the efficiency and selectivity of the product depend on the type of catalyst. The most crucial problem is a low quantum yield in the photocatalysis process due to electron and positive hole recombination, hence produce only limited reduction species on catalyst surface.

In order to solve low quantum yield and selectivity problems many researchers attempted to modify TiO₂ by doping it with metal impurities. Many transition metal ion dopants have been demonstrated to enhance the rate of photocatalytic oxidation or reduction by changing the dynamics of electron-hole recombination and interfacial charge transfer [9]. In CO₂ photoreduction, Yamashita et al. [3] reported that addition of copper (II) to the TiO₂ matrix could improve the efficiency and selectivity to produce methanol. It was suggested however that copper in the first oxidation state (Cu') might play a significant role in the formation of methanol [3]. Tseng et al. has found that the formation of methanol much more effective on Cu-titania catalysts, that the copper species were predominantly Cu₂O [1]. Other investigators also noted that addition of copper could improve the photocatalytic activity in CO₂ reduction, without exactly noticing in which species they are present [2,10]. It is therefore important to investigate how and which copper species (Cu^{0}, Cu^{I}, or Cu^{II}) actually has a significant contribution in improving the CO₂ photocatalytic reduction over TiO₂ matrix.
In the present study, we will report and discuss our finding that each copper species play a different role. A modified preparation method has been conducted to synthesize a Cu/TiO₂, Cu₂O/TiO₂, and CuO/TiO₂ catalysts. The measurement of these catalysts by means of XRD, UV-vis diffuse reflectance spectroscopy (DRS), SEM/EDAX/Mapping and BET have been carried out to investigate the characteristics of the catalysts. Special attention has been focused on the relationship between the copper species and the photoefficiency of methanol formation. Therefore temperature dependence of the reaction and in situ FTIR study were also performed to clarify the kinetic and mechanistic aspects, respectively.

**MATERIALS AND METHODS**

**Preparation and Characterization of Catalysts:** Copper/TiO₂ catalysts were prepared by impregnating TiO₂ Degussa-P25 with copper nitrate solution to give copper content that varies between 0.5 and 10 wt%. The improved-impregnation method was done by conventional impregnation process that followed by sonication of the impregnated TiO₂ slurry, stirring of the slurry at 368 K, drying followed by calcination and reduction-oxidation treatment to form Cu/TiO₂ and Cu₂O/TiO₂ catalysts. Product of the atmospheric calcination step (773 K, 0.5 h) is predicted to be in the form of CuO/TiO₂. Reduction step under 100 ml/min of 20%H₂-He mixture at 573 K for 3 h was completed to make a metallic copper form (Cu/TiO₂). The final step of oxidation was performed under 100 ml/min of N₂O at 353 K for 3 h to get Cu₂O/TiO₂.

The specific surface area of catalysts was measured by multipoint BET of N₂ adsorption at 77 K in a Quantachrome Autosorb-6. Prior to BET analysis, the catalysts were degassed at 423 K under vacuum for 10-12 h. The crystalline phase of the catalysts was identified using powder X-ray diffraction (XRD). The XRD patterns were obtained from a PHILIPS PW 1710 using Ni-filtered Cu Kα radiation (λ=0.154184 nm) that operated quartz reactor cell equipped with ZnSe windows and a copper anode. The 2θ scanning range was 23°-63° connected to a conventional vacuum system. The XRD patterns were recorded for 20-25 min. The 2θ angle was scanned every 5° with a step size of 0.01° and a time per step of 1.0 s. The crystallite sizes of the samples were estimated from FWHM (full-width at half-maximum) of XRD by Scherrer equation. A Cary 2415 UV-vis-NIR spectrophotometer equipped with an integrated sphere was used to record the diffuse reflectance spectra (DRS) of the catalysts. The reflectance spectra of the samples were recorded under ambient condition in the wavelength range of 340-700 nm. A SEM 515 PHILIPS scanning electron microscope (SEM) equipped with EDAX PV9900 PHILIPS energy dispersive X-ray (EDX) and Mapping was conducted at 15 kV and 22 kV to observe the morphology of catalysts, to perform semi-quantitative analysis and to analyze distribution of Cu, respectively.

**Photocatalytic Reaction:** The photocatalytic reduction of CO₂ under UV light was carried out in a reactor system that schematically illustrated in Fig. 1. A flat top cover of the vessel was made of Pyrex and equipped with a gas bubbler and sampling port. The reactor vessel was irradiated, from top of the reactor, by 6 Toki tubular black light UV lamps (@ 10 W), which have the radiation peak about 365 nm. As measured with Black Ray Ultraviolet Meter Model 1225, the UV lamps had total intensity of 2450 µW/cm² in the surface of the catalyst slurry. The reactor was enclosed by an aluminium foil reflector to optimize UV irradiation and to prevent interference from outside light. The reaction temperature was controlled with hot plate magnetic stirrer and fan cooler.

Previous to photo reactions, blank experiments were conducted to ensure that the product formed was due to the photoreduction of CO₂. The blank tests consist of the solution illuminated without the catalyst and a reaction in the dark with the catalyst. Reaction suspensions were prepared to using 300 ml of 1 M KHCO₃ solution and 0.3 g of catalyst powder was added, subsequently followed by sonication for 30 min. Ultra High Purity (UHP) grade CO₂ is bubbled through the reactor for at least 30 min to purge air and to saturate the suspension. The reactor is tightly closed during the reaction and a magnetic stirrer agitated the catalyst-suspended solution at the bottom to prevent sedimentation of the catalyst. The liquid sample was withdrawn every hour from a reactor column. The catalyst-suspended samples were centrifuged to separate catalyst particles from the liquid samples and then analyzed by a GC-FID of Shimadzu 8-ATC equipped with Porapak-Q column.

The in situ IR spectra were measured using an IR quartz reactor cell equipped with ZnSe windows and connected to a conventional vacuum system. The catalysts (27.5 mg) were pressed into a 10 mm diameter disk and then placed in the cell. Prior to photoreactions and IR measurements, the catalysts were heated at 423 K for 2 h and then evacuated at 2x10⁻⁴ torr. UV irradiation of the catalysts in the presence of CO₂ (330 µmol) and gaseous H₂O (1650 µmol) was carried out using the Toki black light UV lamps at 333 K. The spectra on catalyst surface were periodically recorded using a Genesis series FTIR Spectrometer ATI Mattson. Each spectrum was
Fig. 1: The scheme of the photocatalytic reaction system

(1 = Stainless steel vessel, 2 = Gas bubbler, 3 = Control valve, 4 = Sampling Port, 5 = Magnetic stirrer bar, 6 = Inlet fan, 7 = Outlet fan, 8 = UV lamps, 9 = Temperature regulator, 10 = Speed regulator, 11 = Thermometer)

recorded by averaging 150 scans at an instrument resolution of 4 cmG.

RESULTS AND DISCUSSION

Physical Characterization: As shown in Fig. 2, SEM analysis of the catalysts indicated that addition of Cu on TiO₂ can affect the surface morphology of the catalyst; however there is no significant effect on the aggregate sizes. The aggregate sizes of the catalysts were relatively uniform, ranging from 0.4 to 1.0 μm. The EDX/Mapping analysis of Cu on the catalysts demonstrated that Cu uniformly dispersed on TiO₂ surface and further increase of copper leads to the increase of the peak as well as the dot intensities of Cu (inset of Fig. 2).

Figure 3 displays the XRD spectra of the Degussa P25 and copper/TiO₂ of various copper loadings. The peaks on 2θ = 25.3° and 2θ = 27.4° correspond to the main peak of anatase and rutile, respectively. It can be confirmed that for all copper modified TiO₂ samples, only the characteristic peaks corresponding to P25 were found (which consisted of 79% anatase and 21% rutile). Small diffraction peaks of CuO appeared near 2θ = 35.6° and 2θ = 38.8° on the high CuO loading samples ($3\%$) and the peak intensities increase for higher CuO loading. The crystalline size of anatase and rutile were relatively uniform, ranging from 19 to 22 nm and 23 to 31 nm, respectively. Whereas the copper species clusters grew up from 16 to 25 nm with increasing of the loading from 3 to 10 wt%, respectively. No CuO peaks were observed on the low CuO loading samples is due to either the slight amount of copper loading or very small CuO particle size that results in high dispersion of the dopant species. In addition, the disappearance of the CuO peaks after reduction of the samples (Fig. 3-d') verified that CuO has been successfully reduced to other species.

The dispersion capacity of CuO on TiO₂ (anatase) should be equal to the number of the surface octahedral vacant sites available in the preferentially exposed (001) plane of anatase, i.e., 6.98 Cu²⁺ nm² [11]. When the CuO loading exceeds its dispersion capacity, all the vacant sites have been occupied and then the existence of crystalline CuO can be detected by XRD. Plotted in inset of Fig. 3 is the ratio of the XRD peak intensities of CuO and TiO₂ vs. the loading amounts of CuO in the samples, from which the dispersion capacity of CuO on TiO₂ P25 (79% anatase and 21% rutile) is determined to be 2.2 wt.% (4.16 Cu²⁺ nm²). Whereas Xu et al. [11] have investigated that the dispersion capacity of CuO on TiO₂ (pure anatase) is about 6.74 Cu²⁺ nm². The discrepancy of this dispersion capacity may be caused by the different TiO₂ crystalline used as precursor as well as the higher of thermal treatment in this work.

The DRS patterns of TiO₂ P25 and all CuO-loaded samples are shown in Fig. 4. The figure shows that the spectra of copper-titania catalysts are different from unmodified TiO₂ P25. Surface modification of TiO₂ with copper significantly affects the absorption properties of the catalysts. The absorption spectra were obtained by analyzing the reflectance measurement with
Fig. 2: SEM (left) and EDX (right) images of the catalysts. Inset: mapping images of Cu (a = 1%CuO/TiO$_2$; b = 5%CuO/TiO$_2$; c = 10%CuO/TiO$_2$)
Fig. 3: XRD spectra of prepared catalysts: (a) TiO$_2$-P25, (b) 1%CuO/TiO$_2$, (c) 3%CuO/TiO$_2$, (d) 5%CuO/TiO$_2$, (d’) reduced-5%CuO/TiO$_2$, (e) 10%CuO/TiO$_2$.
Inset: plot of quantitative analysis of the XRD data

Fig. 4: DRS patterns of copper-titania catalysts: (a) TiO$_2$ (P25), (b) 0.5%CuO/TiO$_2$, (c) 1%CuO/TiO$_2$, (d) 3%CuO/TiO$_2$, (e) 5%CuO/TiO$_2$, (f) 10%CuO/TiO$_2$.
Inset: plot of bandgap estimation
Kubelka-Munk emission function [9]. It is noticeable that the absorbance of the copper loaded samples increases with increasing Cu content. The absorption edge of TiO$_2$ shifts toward the visible region upon the addition of copper and the baseline in the visible light region is clearly raised.

The bandgap can be estimated by extrapolating the rising segment of the UV spectrum to the abscissa at zero absorption [12]. As shown in inset of Fig. 4, the bandgap of the modified catalysts decreased with increasing of copper loaded. Remarkably, the bandgap is governed by the crystalline structure and the defects in the TiO$_2$ network. Sanchez et al. [12] suggested that small bandgaps were caused by the stoichiometric deficiency of Ti/O ratio.

A specific surface area of unmodified titania catalyst was 54 m$^2$/g, whereas the surface area of copper-loaded titania samples were relatively uniform, ranging from 45 to 50 m$^2$/g. It can be inferred that copper loading on the TiO$_2$ surface did not significantly change the surface area of the catalysts. A small decrease in some catalysts was caused by thermal treatment during the preparation. The surface area values are only somewhat decreased with increase of copper loading. Apparently, the slight change in surface area does not contribute to the reactivity of the catalyst.

**Photocatalytic Activity:** Figure 5 presents the rate of methanol formation on illumination time for various catalysts. Methanol yield was used to evaluate the performance of the catalysts, as it was the major product. Ethanol, propanol, acetone and other hydrocarbons (included oxalate, formate and CH$_3$O) might have been generated, but in small quantities which is too low to be detected. Blank experiments, in the absence of TiO$_2$ as well as in the dark reaction, showed that there was no organic product found for long periods of CO$_2$ photoreduction. Various yields of methanol were obtained in a period depending on the type of catalysts. The maximum methanol yield was achieved by 3%CuO/TiO$_2$ representing that this catalyst had the highest reactivity among all the prepared catalysts.

Figure 6 plots the effect of copper loading on methanol yields, indicating that the methanol yields increased with Cu loading, but then decreased when the Cu loading exceeded 3 wt%. Obviously, more Cu loading can increase methanol yield owing to the amount of active sites. Copper can serve as an electron trapper and prohibits the electron-hole recombination, significantly increasing photoefficiency [1]. However, catalysts with more than 3 wt% Cu loading cannot further increase the methanol yield due to its shading effects which are much higher, consequently reducing the photo exciting capacity of TiO$_2$. Therefore it is estimated, under the experimental conditions of this work, that an optimum amount of copper loading is approximately 3 wt%.

As shown in DRS patterns (Fig. 4), the absorption spectra of low copper-loaded catalysts (#3 3 wt% Cu) were not far different from TiO$_2$ P25 pattern and only shifted to the visible region. It can be supposed, as reported by Anpo et al. [13], that in this case implanted metal ions do not work as electron-hole recombination centers but only work to modify the electronic properties of the catalyst. The band gap values of the copper-loaded catalysts, as revealed in Fig. 4, were smaller than TiO$_2$; consequently the photon energy needed to excite electron would not be too high and furthermore it can increase the photoactivity of TiO$_2$. The dispersion capacity of 2.2 wt% Cu, as evaluated by XRD analysis, indicated that the increasing of CuO above 2.2 wt% might increase the shading effects. However below 3 wt% loading, we found that its shading effects could still be covered by its high ability to trap electrons.

Figure 6 also demonstrates that CuO is the most active dopant comparing with the other species. The formation of methanol was found to be much more effective on Cu$^{2+}$ loaded TiO$_2$ catalyst. One of the considerations in selecting copper oxide as dopant is based on their potential redox position as well as the potential redox value, which is represents their ability to attract electrons. From a thermodynamic perspective, trapping electron by metal ions (Cu$^+$ or Cu$^{2+}$) within the semiconductor photocatalyst is feasible due to the reduction potential of the metal ions is more positive than that of the conduction band edge of TiO$_2$ ( -0.2 V) [9]. The positive potential redox value of Cu$^+$ is higher than that of Cu$^{2+}$ (0.52 vs. 0.34 V); therefore Cu$_2$O dopant should effectively act as an electron trapper to prohibit electron-hole recombination. However, owing to the relatively strong interaction between TiO$_2$ and the dopant particle implanted in the vacant sites of TiO$_2$, the dopant with more positive potential redox exceedingly catches electron from conduction band edge. As a result the dopant-trapped electrons are more difficult to be transformed to the adsorbed species on catalyst surface. The smaller activity of Cu$^+$ and Cu$^{2+}$ species is not associated with lower selectivity by methanol formation due to undetectable of other products in all species of copper-doped Titania catalysts.
On the contrary, Cu\textsuperscript{2+} with the lower potential redox is a more promising dopant species. Due to the reduction of Cu\textsuperscript{2+} is thermodynamically feasible and a Cu\textsuperscript{2+} ion has an unfilled 3d shell, it is valid to assume that electrons can be trapped by CuO\textsuperscript{2+} on the surface of TiO\textsubscript{2} [9]. Figure 7 and equation (1) illustrate the redox cycle of Cu\textsuperscript{2+/2+} that maybe played by CuO dopant on the TiO\textsubscript{2} matrix. Part of the photo-excited electrons on the conduction band (e\textsubscript{CB}) will be trapped by Cu\textsuperscript{2+}, consequently the species of Cu\textsuperscript{2+} can be reduced to Cu\textsuperscript{+} species. Trapped electrons could be consumed via the reduction of H\textsuperscript{+} and/or O\textsuperscript{2-} that present in the system forming radicals of \textit{G\textsubscript{H}} and/or \textit{G\textsubscript{O2}}, respectively. Consequently, the Cu\textsuperscript{+} species could be reoxidized again to Cu\textsuperscript{2+} species. As a result of this sequential cycle, the electron-hole recombination rate could be effectively reduced. In addition, Cu\textsuperscript{2+} species could act as a main adsorption site of CO\textsubscript{2}, for methanol synthesis [14]. Furthermore, the fixation of CO\textsubscript{2} via \textit{G\textsubscript{H}} radicals can lead to methanol formation [7].

\[
\text{Cu}^{2+}\text{O} \xrightarrow{e^-} \text{Cu}^+\text{O} \xrightarrow{\text{O}_2/\text{H}^+} \text{Cu}^{2+}\text{O} \quad (1)
\]

To study the kinetic aspect of the CO\textsubscript{2} photoreduction, the experiment was repeated at different temperatures in the range of 316-373 K. Inset of Fig. 6 shows a typical Arrhenius plot for Degussa P25 and 3%CuO/TiO\textsubscript{2} catalysts. A mean value for apparent activation energy (E\textsubscript{a}) of ca. +26 and +12 kJ/mol was calculated for Degussa P25 and 3%CuO/TiO\textsubscript{2}, respectively. Determination of the E\textsubscript{a} value is quite accurate due to the coefficient correlation R\textsuperscript{2} is in the range of 0.90-0.93. These positive values of E\textsubscript{a} imply that the desorption of products is the rate limiting step in the photosynthetic formation of methanol [15]. In addition, the lower E\textsubscript{a} of 3%CuO/TiO\textsubscript{2} catalyst may indicate a catalytic role of copper species as an active site to provide methanol and hence improving the photocatalytic activity of TiO\textsubscript{2} catalyst. Guan et al. [16] reported that in the photocatalytic reduction of CO\textsubscript{2} under concentrated sunlight, CH\textsubscript{3}OH was catalytically formed over the Cu/ZnO catalyst if it combined with K\textsubscript{2}Ti\textsubscript{3}O\textsubscript{7} photocatalyst.

The in situ IR experiments were conducted to investigate the mechanism aspects of the catalyst. Figure 8 shows the in situ IR spectra of 3%CuO/TiO\textsubscript{2} catalyst during gas phase CO\textsubscript{2} photoreduction as a function of time irradiation. The methanol product was identified by growing up of the absorption bands at 1430-1560 cm\textsubscript{G}, which are assigned to \(*\text{-}(\text{CH})_3\). The absorption bands at 1360 and 2867 cm\textsubscript{G} also correspond to lewis bound of CH\textsubscript{3}OH *(OH) and lewis bound of CH\textsubscript{3}OH *(CH\textsubscript{3}), respectively. The band at 2968 cm\textsubscript{G} corresponds to \lewis{(CH\textsubscript{3})}. Although the bands at 2600-3000 cm\textsubscript{G} are not completely clearly caused by cold to noise, a formate species adsorbed on Ti are observed at 1544, 2873, 2950 cm\textsubscript{G} and the bands at 2845, 2939, 1373 cm\textsubscript{G} are assigned to formate-Cu species. These formate species adsorbed on Cu and Ti can be regarded as an intermediate of the methanol synthesis, as reported by previous researchers [19,20]. Methoxy species, which can be predicted as hydrogenation product from formate species [14,17] detected at 2824 and 2920 cm\textsubscript{G}. However, the methoxy species could not be observed on the surface of undoped-TiO\textsubscript{2} photocatalyst.

Numerous reaction schemes for the photocatalytic reduction of CO\textsubscript{2} on TiO\textsubscript{2} based catalysts have been proposed in different studies [1,5-7]. Although the exact mechanism of methanol production from CO\textsubscript{2} photoreduction is not absolutely clear in our present study, the results obtained in the current analysis show good correspondence with the kinetics as well as in situ IR experiments. The appearance of formate and methoxy intermediates on the CuO/TiO\textsubscript{2} surface, while none on the undoped-TiO\textsubscript{2} catalyst, may indicate that the catalytic role of copper species as an active site is to provide methanol. In the hydrogenation of CO\textsubscript{2}, Kakumoto [14] proposed that the methanol synthesis mainly occurs on Cu\textsuperscript{2+} site which the linear adsorption of CO\textsubscript{2} dominantly take place. The mechanism of methanol synthesis on the CuO/TiO\textsubscript{2} surface is proposed, as shown in the following equations (2 - 5) followed by mechanism scheme in the Fig. 9.

\[
\text{TiO}_2 + 6h^+ + 6\text{H}_2\text{O} \rightarrow 6\text{H}_3\text{O}^+ \quad (2)
\]

\[
\text{H}_2\text{O} + \text{CO}_2 \rightarrow \text{H}^+ + \text{HCO}_3^- \quad (3)
\]

\[
2\text{H}^+ + \text{HCO}_3^- \rightarrow 6\text{H}_2\text{O} + \frac{1}{2}\text{O}_2 + \text{CO}_2 \quad (4)
\]

\[
\text{H}^+ + e^- + 6\text{G}\textsubscript{H} \quad (5)
\]

Photo-excited electrons (e\textsuperscript{-}) and positive hole (h\textsuperscript{+}) are generated when incident photons are absorbed in the surface of CuO/TiO\textsubscript{2} catalyst. The holes first react with water and/or H\textsubscript{2}CO\textsubscript{3}, resulting in producing hydrogen ions (H\textsuperscript{+}) and oxygen. Moreover generated electrons may be consumed via the reduction of H\textsuperscript{+} forming radicals of \textit{G\textsubscript{H}}. At the same time, CO\textsubscript{2} is adsorbed on Cu\textsuperscript{2+} site linearily. Furthermore, it is suggested that the sequential interaction of \textit{G\textsubscript{H}} radicals and H\textsuperscript{+} with the adsorbed state
Fig. 5: Time dependence on the methanol yields of various catalysts at 333 K and UV irradiation.

Fig. 6: Effect of copper loading amount on methanol yield under 333 K and 6 h UV irradiation. Inset: Arrhenius plot for TiO₂ and 3%CuO/TiO₂ catalysts.

Fig. 7: Schematic of the redox cycle of Cu²⁺/Cu⁺ system in the photoreduction of CO₂.
Fig. 8: *In situ* IR spectra of 3% CuO/TiO$_2$ catalyst during gas phase CO$_2$ photoreduction (Mol ratio of H$_2$O/CO$_2$ = 5, CO$_2$ = 330 µmol, T = 333 K)

Fig. 9: Scheme of CO$_2$ photoreduction mechanism on CuO/TiO$_2$ catalyst
of CO₂ on Cu⁺ surfaces lead to the formation of methanol via the formate and methoxy intermediates.

CONCLUSIONS

The photocatalytic reduction of CO₂ with H₂O at the liquid/solid interface of copper-doped titania photocatalysts have been performed, giving methanol as a main product. Experimental results demonstrated that the photocatalysts with a various dopant species (CuO, Cu₂O, or Cu⁵⁺) can be synthesized by an improved-impregnation method via the reduction-oxidation steps. The optimal Cu-loaded TiO₂ (in this study, 3 wt %) has shown to be a highly efficient photocatalyst for CO₂ photoreduction. It has been revealed that CuO is the dopant species that actually has a significant contribution in improving the CO₂ photoreduction over TiO₂ matrix. The apparent activation energy (Eₐ) for 3%CuO/TiO₂, in the working temperature (316-373 K), is lower than that for Degussa P25. This suggests that CuO dopant species plays a role as a catalytic site in enhancing the production of methanol. in situ FTIR experiment revealed that the methanol was also catalytically synthesized on the CuO/TiO₂ surface by producing formate and methoxy intermediates. Therefore, the considerable improvement of CO₂ reduction photoefficiency by CuO dopant is proposed due to the double effects of the dopant: i.e. as an electron trapper and a catalytic active site.

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