CO₂ Splitting by H₂O to CO and O₂ under UV Light in TiMCM-41 Silicate Sieve

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The 266-nm light-induced reaction of CO_2 and H_2O gas mixtures (including isotopic modifications $^{13}CO_2$, $C^{18}O_2$, and D_2O) in framework TiMCM-41 silicate sieve was monitored by in situ FT-IR spectroscopy at room temperature. Carbon monoxide gas was observed as the sole product by infrared, and the growth depended linearly on the photolysis laser power. H_2O was confirmed as the stoichiometric electron donor. The work identifies CO as the single-photon, 2-electron-transfer product of CO_2 at framework CO_2 at framework CO_2 acting as an electron donor, which has not been established before. CO_2 was detected as coproduct by mass spectrometric analysis of the photolysis gas mixture. These results are explained by single CO_2 by CO_2 by CO_3 by CO_3 and surface CO_3 and surface CO_3 by CO_3 and surface CO_3 and surface CO_3 by CO_3 and surface CO_3 and surface CO_3 by CO_3 and CO_3 and surface CO_3 and surface CO_3 by CO_3 and CO_3 are CO_3 and CO_3 and

1. Introduction

The photoreduction of CO₂ by H₂O is one of the most important chemical reactions for solar energy-to-fuel conversion. First reports on the reaction by a heterogeneous photochemical approach were by Hemminger et al. and by Inoue et al. using TiO2, SrTiO3, or SiC semiconductor powders or single crystals.^{1,2} Band gap excitation by UV light led to the formation of methane and methanol as products. Most semiconductor studies were conducted with aqueous suspensions of colloidal particles of TiO2, ZnS, ZnSe, CdSe, etc., often surface-modified to enhance efficiency, selectivity, or wavelength response.^{3,4} Reduction is initiated by transfer of photogenerated conduction band electrons to surface-adsorbed CO2 or carbonate. For materials that are stable under use (metal oxides), excitation wavelengths are limited to the UV region. On the other hand, visible light-induced reduction of CO₂ to CO or formic acid is achieved with low band gap semiconductor particles such as ZnSe, ZnS, or CdS, but these semiconductors require sacrificial reductants to suppress irreversible oxidation of the material. Similarly, visible light-activation of CO₂ in the presence of transition metal colloids requires sacrificial organic donors.⁵

Substantially improved yields of CH₃OH and CH₄ from CO₂ and H₂O under UV light were found by Anpo and co-workers over the past several years by engaging isolated tetrahedral Ti centers of micro- or mesoporous silicate sieves instead of dense phase TiO₂ materials.^{6,7} Molecular sieves used are microporous Ti silicalite (TS-1),⁸ zeolite Ti- β ,⁹ or mesoporous framework substituted Ti silicates such as TiMCM-41.8,10 The product distribution between methanol and methane was influenced by the hydrophilic character of the solid. Overall yields were found to be higher for mesoporous than microporous silicate hosts. Development of framework Ti-containing mesoporous silicate films that are optically transparent (as opposed to the lightscattering pressed wafers of crystallites) has allowed the determination of the quantum efficiency, which is around 0.3% (300 nm). 11,12 UV photolysis of CO₂ and H₂O gas mixtures loaded into the sieve was typically conducted at moderately elevated temperature (around 325 K) and monitored by GC analysis of the products. Methanol and methane were the main products, along with trace amounts of CO, O2, C2H4, and

C₂H₆.^{6,7} Mechanistic insights were sought by EPR spectroscopy of TiO₂ samples irradiated at 77 K, resulting in the detection of H atoms and carbon radicals (elemental C and CH₃).^{13,14} It was proposed that CO₂ reduction and H₂O splitting proceed competitively at the LMCT-excited Ti-O centers: CO₂ is reduced to CO, and subsequently to C radicals while H₂O photodecomposes to H and OH radicals. Reaction of H and OH radicals with carbon species is thought to yield CH₃OH and CH₄.^{8,13,14}

Understanding of the mechanism of UV light-induced reduction of CO2 by H2O in Ti silicate sieves under reaction conditions would greatly aid in the design of photoreactors that accomplish the chemistry at longer wavelengths. Of particular interest is the identification of the individual reaction steps leading to the final products, photon by photon. Equally important is knowledge of the number of quanta needed to reach CH₃OH (6-electron transfer) or CH₄ (8 electrons). Moreover, possible distructive cross reactions of intermediates or final products need to be identified since they invariably limit the quantum efficiency. In our first attempt to address these questions, we report here an in situ FT-IR and mass spectrometric study of 266-nm-induced CO₂ reduction by H₂O in framework TiMCM-41 silicate sieve at room temperature. The work led to the identification of the single-photon 2-electron products under reaction conditions.

2. Experimental Section

TiMCM-41 sieve was prepared following a modified method for MCM-41 synthesis, ¹⁵ but now including a Ti precursor in the synthesis gel. Briefly, 2.2 g of CTAB template (cetyltrimethylammonium bromide, Aldrich) were dissolved in 52 mL of water at 40 °C. Ammonium hydroxide (Aldrich, 26 mL) was then added under stirring. Addition of 10 mL of tetraethoxysilane (Fluka, puriss.) and 0.03 g of Ti tetrabutoxide to the solution was followed by continued stirring for 3 h at room temperature. The gel was transferred to a Teflon-lined autoclave and held at 110 °C for 48 h. Filtration and washing with distilled H₂O gave the product. Template removal was typically conducted by heating at 300 °C for 2 h followed by calcination under O₂ flow at 550 °C for 12 h. Template removal in some

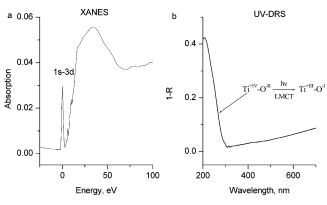


Figure 1. Spectroscopic characterization of framework TiMCM-41. (a) XANES spectrum of Ti K-edge. The A_1 - T_2 preedge peak position (4968 eV) is taken as the origin of the energy scale. (b) UV—vis diffuse reflectance spectrum of calcined TiMCM-41 in a vacuum. R = reflectance, BaSO₄ as the reference.

cases was also done by the ion exchange method, using 0.2 M (NH₄)₂SO₄ in an ethanol—H₂O 1:1 solution. ¹⁶ As-synthesized TiMCM-41 (100 mg) was dispersed in 30 mL of solution and stirred at 65 °C for 30 min, filtered while hot, and washed with cold ethanol. The procedure was repeated 3 times and followed by additional washing with water and ethanol to remove excess ammonium sulfate. The product was dried at 100 °C for 12 h before use. Chemical analysis indicated at Ti/Si ratio of 0.020 \pm 0.001. Powder XRD measurements (Siemens model D500 Cu K α) of the sieve showed the same well-known Bragg peaks of neat MCM-41. ¹⁷ No extra peak or phase was detected. The parent MCM-41 material was prepared by the same procedure as TiMCM-41 without the Ti source.

XANES spectra of the Ti K-edge of pressed wafers of TiMCM-41 recorded at beamline 9.3.1 at the Advanced Light Source showed a sharp, intense preedge peak at 4968 eV (for experimental details of the X-ray absorption measurement, see ref 18). The band, shown in Figure 1a, is attributed to the A₁-T₂ absorption of tetrahedrally coordinated Ti centers. Its height relative to that of the K-edge is a measure of the fraction of Ti in tetrahedral coordination.¹⁹ Comparison with literature XANES spectra of TiMCM-41confirmed the high yield of tetrahedrally substituted Ti centers.²⁰ Furthermore, the UV diffuse reflectance spectra of the material (Shimadzu model UV-2100 spectrometer equipped with an integrating sphere model ISR-260) exhibit an absorption with an onset at 330 nm (Figure 1b). The short wavelength onset of the band shows that it originates from the LMCT transition of tetrahedrally coordinated Ti, in agreement with literature spectra.²¹

For photochemical experiments, 10 mg of TiMCM-41 powder was pressed into a self-supporting wafer. The wafer was mounted in a miniature infrared vacuum cell equipped with CaF₂ windows.²² Dehydration of the sieve was accomplished by room temperature evacuation (turbomolecular pump Varian model V-70) prior to a 3-fold exchange with D₂O vapor (12 Torr). Reactants were loaded into the molecular sieve from the gas phase through a vacuum manifold. Chemistry was monitored in situ by FT-IR spectroscopy at 0.25-cm⁻¹ resolution (Bruker models IFS88 or IFS66V equipped with LN2 cooled MCT detectors model Kolmar KMPV8-1-J2 with an 8 μ m band gap). Photolysis was conducted at room temperature, using the 266nm emission of a pulsed Nd:Yag laser at 10 Hz (Quanta Ray model DCR2A with GCR3 upgrade). For monitoring of O₂ formation, the gas phase of the infrared cell was probed before and after photolysis by a quadrupole mass spectrometer Pfeiffer model Omnistar 422.

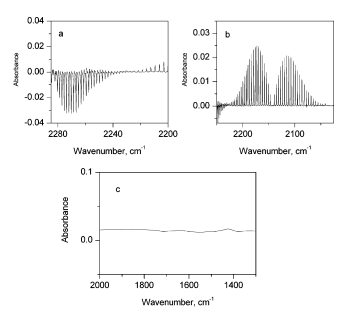


Figure 2. FT-IR difference spectrum upon 266-nm irradiation of TiMCM-41 loaded with 705 Torr of CO_2 and 10 Torr of D_2O for 230 min at 100 mW cm⁻². The negative ro-vibrational bands in parts a and b are due to depletion of $^{13}CO_2$ (natural abundance in $^{12}CO_2$).

CO₂ (Matheson, 99.995%), ¹³CO₂ (ISOTEC, 99% ¹³C), C¹⁸O₂ (ICON, 98.7% ¹⁸O), CO (Matheson, 99.99%), and ¹³CO (Matheson, 99% ¹³C) were used as received. Triply distilled H₂O and D₂O (Aldrich, 99.96%D) were degassed by 4 freeze-pump—thaw cycles before use.

3. Results

Loading of 12 Torr of H₂O (D₂O) into the miniature infrared cell containing a TiMCM-41 wafer gave rise to a very broad water stretching absorption with a maximum at 3300 cm⁻¹ (2500 cm⁻¹) and a bending mode at 1630 cm⁻¹. D₂O was chosen in many experiments to keep the 1600-1700-cm⁻¹ range free of reactant absorption. Possible CO2 reduction products such as formic acid or formate ion²³ would be difficult to detect in the presence of H₂O with its strong bending mode absorption. Subsequent addition of 700 to 750 Torr of ¹²CO₂, ¹³CO₂, or ¹²C¹⁸O₂ resulted in gas-phase infrared bands reported previously.²³ No peaks due to carbon dioxide adsorbed on the sieve were observed except for a small band at 1381 cm⁻¹ (¹³CO₂, 1366 cm⁻¹; $C^{18}O_2$, 1334 cm⁻¹), assigned to the ν_1 symmetric stretch of CO₂.²⁴ While infrared-forbidden for the gas-phase molecule, the mode is induced by interaction with the nanopore environment in the case of the adsorbed CO₂.

When irradiating the loaded TiMCM-41 sieve with 266-nm light at room temperature, growth of gas-phase CO was detected by its ro-vibrational bands in the 2200-2000-cm⁻¹ region. At room temperature, CO generated in the silicate pores desorbs instantly into the gas phase. Figure 2 shows the infrared difference spectrum upon irradiation of a cell containing 705 Torr of CO_2 and 10 Torr of D_2O for 230 min at 100 mW cm⁻². Aside from CO (trace b), the only additional growth was observed in the fingerprint region (trace c) at 1420 cm⁻¹ and is due to HDO, presumably the result of continued slow exchange of D₂O with residual Si-OH groups of the pore surface. No other products were detected by infrared in the 4000-1300cm⁻¹ region. Specifically, no bands grew in the 2300-2200cm⁻¹ range (trace a) where C-D stretch modes of possible methanol²⁵ or methane²⁶ products would absorb. Likewise, no absorbance growth was noticed in the carbonyl stretching region (Figure 2c) where formic acid (DCO₂D, 1695 cm⁻¹),²³ formate

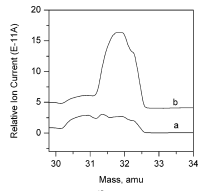


Figure 3. Mass spectrum of a $^{13}\text{CO}_2$ (740 Torr) + H₂O (12 Torr) mixture contained in the miniature IR cell (a) before photolysis and (b) after 266-nm irradiation for 300 min at 160 mW cm⁻². The signal is the average of four photolysis experiments, each conducted with a fresh TiMCM-41 wafer.

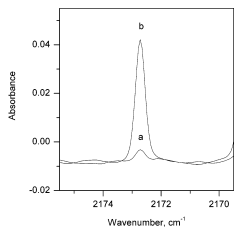


Figure 4. H_2O dependence of CO_2 photoreduction. (a) CO absorbance growth at 2173 cm⁻¹ upon 266-nm irradiation of TiMCM-41 loaded with 1 atm of CO_2 for 75 min at 194 mW cm⁻². (b) Same photolysis experiment as in part a with TiMCM-41 loaded with 1 atm of CO_2 and 12 Torr of H_2O .

 $(DCO_2^-, 1565 \text{ cm}^{-1})$, 23 or carbonate $(CO_3^{2-}, 1440 \text{ cm}^{-1})$; HCO₂⁻, 1600 cm⁻¹)²⁷ are known to absorb. In a series of experiments with ${}^{13}\text{CO}_2$ (740 Torr) + D₂O (12 Torr) mixtures, the gas content of the infrared cell (3.4 cm³ volume) was transferred to a quadrupole mass spectrometer for analysis before and after photolysis. Figure 3 shows the average signal observed at mass 32 in four experiments, each conducted with a new TiMCM-41 wafer. Trace a is the spectrum before photolysis; the small signal at mass 32 is due to O₂ generated by ¹³CO₂ fragmentation in the mass spectrometer. Figure 3b shows that O_2 is produced by photolysis of ${}^{13}CO_2 + H_2O$ at 266 nm. While well above background for each run, the signal fluctuated from experiment to experiment by as much as a factor of 4. These fluctuations can be understood on the basis of the proposed mechanism of O2 generation, namely dismutation of surface OH radicals, discussed in Section 4. We conclude that gas-phase CO and O₂ are the only products formed upon 266-nm photolysis (<200 mW cm⁻², 5 h) of CO₂ and H₂O in TiMCM-41 at room temperature.

Carbon monoxide is only produced when water is present as donor. Figure 4b shows the infrared absorbance growth of a single ro-vibrational line of CO at 2173 cm⁻¹ upon irradiation of a $CO_2 + H_2O$ mixture in TiMCM-41 for 75 min at 194 mW cm⁻². Trace a of Figure 4 is the spectrum observed when repeating the same photolysis experiment without added H_2O .

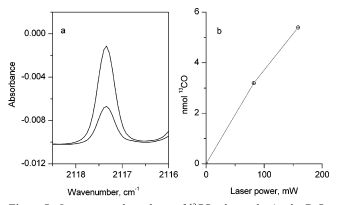


Figure 5. Laser power dependence of ¹³CO₂ photoreduction by D₂O. (a) Spectra of the ¹³CO band at 2117 cm⁻¹ recorded after 30 and 80 min of photolysis. (b) Laser power dependence of ¹³CO absorbance growth. Each data point represents the average of 8 measurements.

We can only speculate at this point about the possible source of the small CO growth in Figure 4a, which occurs only during the initial 30 min of irradiation. One possibility is the residual H₂O present in the sieve. It would imply that water molecules regenerated upon dismutation of the primary products (see Figure 6) are no longer accessible for continued reaction at the redox sites, which is reasonable to assume because of the presence of 1 atm of CO₂.

Infrared monitoring of ${}^{13}\text{CO}_2 + \text{H}_2\text{O}$ photolysis revealed the formation of some ¹²CO gas along with the ¹³CO product, with the ¹²CO growth fluctuating erratically between experiments. This points to the presence of small amounts of carboneous residues that react with oxidizing intermediates formed during the photochemical reaction. Such carboneous residues are known to be difficult to remove completely from the high-surface-area mesoporous silicates.^{28,29} The origin of ¹²CO was confirmed experimentally by our finding that the ¹²CO yield decreased by a factor of 2 when conducting the experiment with a C18O2 + $\rm H_2^{16}O$ mixture instead of the $^{13}C^{16}O_2 + \rm H_2^{16}O$ sample. The result demonstrates that half of the O atoms of the ¹²CO impurity produced by the photochemical reaction originate from CO₂, the other half from H₂O. Therefore, the carboneous residues in the sieve act as a trap for transient oxygen species that otherwise would react to form O_2 .

To determine whether single or multiple photons are required for the CO production, we examined the laser power dependence of the ¹³CO absorbance in ¹³CO₂ + D₂O experiments. Figure 5a shows the growth of the ro-vibrational line of ¹³CO at 2117 cm⁻¹ that was used for the analysis. As can be seen from the bands measured after consecutive photolysis periods, the fwhm is constant in the growth range used for the analysis, permitting the use of peak absorbances. Experiments were conducted by measuring the growth of the 2117-cm⁻¹ peak for 16 consecutive photolysis periods of 5 min duration, alternating between 80 and 160 mW cm⁻² laser power. Absorbance growth data for each of the two power levels were averaged. Calibration of the 2117-cm⁻¹ band intensity in terms of ¹³CO pressure by using an authentic ¹³CO sample allowed us to determine the quantity of ¹³CO produced in the IR cell. As can be seen from Figure 5b, the carbon monoxide growth depends linearly on the photolysis laser intensity. The slightly less than doubling of the growth at 160 mW cm⁻² is consistent with the additional thermal heating by a few degrees caused by the increased laser power, resulting in slightly less adsorption of reactants. Since saturation effects can be ruled out at the modest photolysis light intensities used, the linear dependence implies that CO is produced by a single photon process.

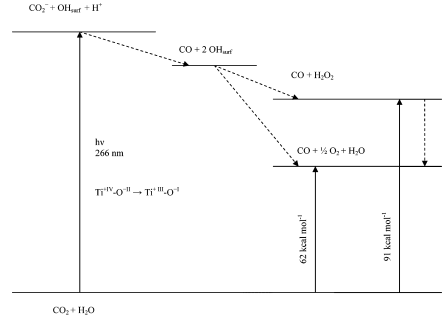


Figure 6. Proposed mechanism for UV light-induced CO_2 reduction by H_2O at tetrahedral framework Ti centers. Energies are accurate except for surface-stabilized OH radicals (see text).

4. Discussion and Conclusions

The essential results of the spectroscopic study of the photolysis of carbon dioxide and water in TiMCM-41 sieve are the establishment of CO as single-photon 2-electron-transfer product, and O_2 as coproduct under reaction conditions at room temperature. Water is confirmed as the stoichiometric electron donor. Both CO and O_2 were already detected previously by Anpo and co-workers as trace products by GC measurements.^{6,7} The importance of the work reported here is the finding that single-photon excitation of the Ti–O LMCT chromophore results in splitting of CO_2 to CO with H_2O acting as the direct electron donor, which has not been established before. The oxidizing fragments generated by the H_2O oxidation and CO_2 dissociation react to produce O_2 .

On the basis of these observations, the mechanism shown in Figure 6 is proposed. Excitation of the Ti-O LMCT transition of Ti centers leads to transient Ti+III and a hole on a framework oxygen. Electron transfer from Ti^{+III} to CO₂ splits the molecule into CO and O-. The latter is spontaneously protonated by a Si-OH group, or H⁺ co-generated upon H₂O oxidation to yield a surface OH radical. Another such surface OH radical is formed as a result of the concurrent H₂O oxidation by the framework oxygen hole. The OH radicals either combine to yield H_2O_2 or dismutate to give O2 and H2O. In this UV photolysis experiment it is not possible to determine whether hydrogen peroxide is formed because, according to our study of H₂O₂ interacting with Ti silicate, it can react with Ti centers to yield TiOOH moieties. The latter photodissociates to H₂O and oxygen at wavelengths shorter than 500 nm.³⁰ Figure 6 indicates the free energies associated with the formation of the stable products. The only speculative energy is that of the surface OH radicals. The free energy of the reaction $CO_2 + H_2O \rightarrow CO + 2$ OH, with all species in the gas phase, is +139.7 kcal mol⁻¹.³¹ The maximum energy available in a single-photon process with 266-nm quanta is 107 kcal mol⁻¹. This implies that the OH species must be stabilized by interaction with the silica pore surface by at least 16 kcal mol⁻¹ per radical, which is plausible. In light of the long lifetime of even very small radicals in a room temperature molecular sieve due to random walk (hundreds of microseconds for HCO in zeolite Y),³² it is not surprising that, even at a very low concentration, carboneous residues offer a significant trapping mechanism for OH radicals.

It is interesting to note the difference in the 2-electron-transfer product of CO₂ reduction on a Ti silicate sieve depending on whether H₂O or CH₃OH is used as donor. We have previously found that CO2 photoreduction in Ti silicalite when using methanol as donor generates HCO₂H, not CO, as the primary reduction product.²³ This is most probably due to the fact that one-electron oxidation of CH₃OH by the hole on framework O produces the CH₂OH radical, a H atom donor that can reduce transient CO₂⁻ to HCO₂H. By contrast, no H atom donor is generated upon H₂O oxidation that would quench CO₂⁻, resulting in spontaneous splitting of the transient. In fact, the lack of accumulation of highly reactive reducing radicals constitutes an advantage of photochemical CO2 splitting by H2O over other photosynthetic reactions because back reaction of reduced and oxidized transient radicals does not occur. The reduced species formed by CO₂ splitting (CO) desorbs from the reactor, leaving behind the same oxidizing species (OH) generated by photooxidation of the donor.

With CO_2 splitting identified as the key activation step, redox centers can be explored that may afford the reduction of CO_2 by H_2O with photons at longer wavelengths.

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