Photocatalytic System using Iron-Based Catalysts: Effect of External Ions on CO₂-to-CO Conversion

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Summary: Herein, we report four self-assembly Iron(II/III) complex consisting triethanolamine (TEOA) as ligand and an Fe^{III}(TPA)Cl₃ (TPA= 3-(2-pyridine methyl) amine) as contrast catalyst, for photochemical reduction of CO₂ to CO under visible light. The photocatalytic systems were assembled by [Ru(bpy)₃]Cl₂ as photosensitizer, and 1,3-dimethyl-2-phenyl-2,3-dihydro-1H-benzo-[d]-imidazole (BIH) as electron donor. Experiments are conducted to study the relationship between CO production and the external ions. Different external counterions are provided to assist the discussion of the influence of these external ions on catalysis. Density functional theory calculations confirmed that the external ions have an effect on the CO yield. Moreover, a mechanism involving proton-coupled electron transfer is proposed.

Key words: Photocatalytic, CO₂-to-CO conversion, External ions, Iron-Based Catalysts, Density functional theory.

Introduction

Visible-light-driven reduction of CO₂ to CO has been of considerable interest in the area of photochemistry for many years. Since Lehn et al. pioneering work in 1986, in which the photocatalytic reduction of CO₂ using Co polypyridine complexes in a homogeneous system [1], the development of low-cost, highly efficient and stable photocatalysts remains a constant challenge. Recent studies have mainly focused on the development of efficient and inexpensive photocatalytic systems. Molecular catalytic systems containing Fe complexes as catalysts have exhibited its potential for the photocatalytic reduction of CO₂. However, examination of the influence of external ions toward CO production using non-noble metal catalysts, such as Fe-based catalyst [2-6], is scarce and worthy further exploration. P. Neta et al. [7] and our group [8] reported [Fe(TEOA)₂]Cl₂ as catalysts and only a low efficiency of CO₂-CO conversion or hydrogen evolution was obtained. Herein, we utilized [Ru] ($[Ru(bpy)_3]Cl_2$) as PSs and BIH (1,3-dimethyl-2-phenyl-2,3-dihydro-1Hbenzo-[d]-im idazole) as sacrificial reductant for photocatalytic reduction CO2 to CO. In this study, instead of focusing on the preparation of new photocatalysts, we concentrated on studying the simple Fe-based catalysts for the production of CO by tuning the external ions and valence state of Fe.

Experimental and discussion

examined photocatalytic employed Fe complexes with two kinds valence states, Fe(III) and Fe(II), and three anions Cl⁻, Br⁻ and SO₄², Fe^{III}(TPA)Cl₃. (TPA= 3-(2-pyridine methyl) amine) was prepared as a contrast catalyst. The following complexes were prepared and utilized as catalysts: Fe^{III}(TEOA)₂Cl₃; Fe^{II}(TEOA)₂Cl₂; Fe^{III}(TEOA)₂Br₃; Fe^{II}(TEOA)₂SO₄; Fe^{III}(TPA)Cl

a HO
$$CH_2$$
 H_2C
 H_2C

Structures of five catalysts and BIH: a. TEOA as the ligand, X=2 or 3, the anions are Cl⁻, Br⁻ or SO₄²⁻, Fig. 1: respectively.

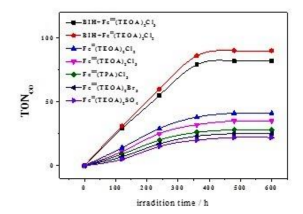
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Comparative experiments were studied using a variety of iron salts (FeCl₃, FeCl₂·2H₂O, FeBr₃ and FeSO₄·7H₂O). All experiments were performed in homogeneous solutions. **DMF** (N. Dimethylformamide) was used as solvent and mixed with TEOA in ratio of 1/5 (v/v, Fig. S1). In each of these experiments, 0.2 mM [Ru] was used as the photosensitizer. Since TEOA is a weak base (pK_a=7.8, in water), Fe ions are easily formed to polymeric hydroxides upon exposure to air. Accordingly, the reaction solutions were deoxygenated with argon before the addition of catalysts. The solution's pH (from 7.8 to 4.1) was adjusted via addition of CH₃COOH, as a consequence, the concentration of Fe polymeric hydroxides decreased, however, a decrease in catalytic efficiency occurred (Fig. S2). TEOA played multiple roles in the reaction process, acting as an electron donor [9] and CO2 adsorbent [10], as well as a ligand [11-13]. TEOA coordinates with Fe³⁺ and Fe²⁺ in solutions and generates the TEOA complexes (Fe^{III}(TEOA)₂Cl₃, Fe^{II}(TEOA)₂Cl₂, Fe^{III}(TEOA)₂Br₃, and Fe^{II}(TEOA)₂SO₄), respectively. The solutions to be irradiated were firstly purged with CO₂ until saturated. The temperature of all solutions was controlled at room temperature using circulatory alcohol as the cooling liquid.

Fig 2 shows the photocatalytic production of CO with different Fe complexes as catalysts under visible light (410 $\leq \lambda \leq$ 750 nm) irradiation. In the case of Fe^{II}(TEOA)₂Cl₂, its turnover number (TON) of 32 (109 µmol) for CO production was greater than the other studied complexes (Fe^{III}(TEOA)₂Cl₃ 28, Fe^{III}(TEOA)₂Br₃ 15.5 and Fe^{II}(TEOA)₂SO₄ 12.5) and $Fe^{III}(TPA)Cl_3$ (TPA = 3(2-pyridine methyl) amine, $TON_{CO} = 18.5$). Interestingly, when BIH (1,3dimethyl-2-phenylbenzimidazoline) was added to the photocatalytic system using Fe^{II}(TEOA)₂Cl₂ as catalyst, TON of CO production increased 3-fold to 89 (303 µmol). The reason for this phenomenon is due to the BIH being a stronger electron-donor than TEOA [14]. Furthermore, a large excess of TEOA restrained the dissociation of TEOA ligand from the metal center. promoting the stability of the complex catalysts.

After 10 h of irradiation and reached the point where CO₂ no longer produces CO, for photocatalytic systems with Fe^{II}(TEOA)₂Cl₂ and Fe^{III}(TEOA)₂Cl₃ as catalysts, respectively, the same amount of photosensitizer [Ru] were reloaded, and in the comparative experiments, no [Ru] was added to the photocatalytic systems, only the Fe^{III}(TEOA)₂Br₃ and Fe^{II}(TEOA)₂SO₄ as catalysts are reloaded, respectively. Under CO₂ atmosphere, the catalysts Fe^{II}(TEOA)₂Cl₂ and Fe^{III}(TEOA)₂Cl₃ were added again into the same photocatalytic system with another

10 h of irradiation (Table S1), TON_{CO} (turnover of number) were 31 and 27.1 once again, respectively. Furthermore, when the other photocatalysts where subjected to the same experiment, CO production were not detected. This result indicates that catalysts Fe^{II}(TEOA)₂Cl₂ and Fe^{III}(TEOA)₂Cl₃ are durable during the photocatalytic process, and that the decomposition of [Ru], rather than catalysts, may be the major constraint limiting durability of the system. Hence, we can Fig out that the Fe^{II}(TEOA)₂Cl₂ and Fe^{III}(TEOA)₂Cl₃ could acted as photocatalysts for reduction of CO₂ with [Ru], and Fe^{II}(TEOA)₂Cl₂ was the most efficient one among these five complexes studied.



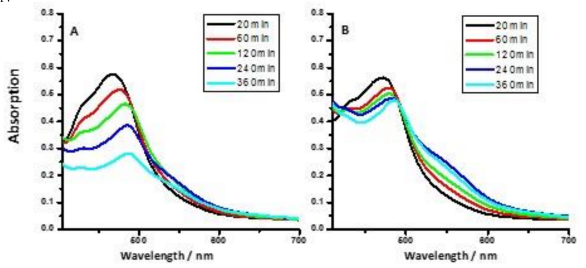
Photocatalytic formation of CO formed from Fig. 2: CO₂ as a function irradiation time: solution [TEOA/DMF = 1/5(v/v)] that contained [Ru](0.2 mM) and one of Fe complexes (preprepared or self-assembled, 1.9 mM) or BIH (29 mM) and all of them saturated by CO₂. Solution were irradiated using a Xe lamp through filters and giving light within the scope of $410 \le \lambda \le 750$ nm.

Isotopic labeling experiment using ¹³CO₂ as the substrate confirmed that the CO product originates from the photocatalytic conversion of CO₂ (Fig. S3).

The number of incident photons, measured using a classical iron ferrioxalate (K₃Fe(C₂O₄)₃) chemical actinometer, was 2.01×10^{24} photons per hour. According to quantum yield calculations [15] and using known parameters [16], the quantum yield of CO formation is $\Phi_{co} = 0.00072$ %.

Fig 3 shows the electronic absorption spectra of [Ru] during photolysis in TEOA and DMF mixed solution, which was performed by UV-Vis spectrophotometry. The spectra shows the metal-toligand $d\pi \rightarrow \pi^*$ charge transition about 455 nm [17]. With the growth of the photolysis time, [Ru] decomposes rapidly in the absence of FeCl₃ (Fig A in Fig 3). However, FeCl₃ added to the system, the decomposition of [Ru] was sluggish (Fig B in Fig 3). As shown in the Fig 3, [Ru] is excited and reductively quenched by TEOA, whose concentration is at least 200 times higher than [Ru](0.2 mM [Ru] vs 50 mM TEOA at minimum). Since there is no CO generated when the system is absent of TEOA. We assume that BIH is only a more superior electron donor than TEOA, while TEOA is both a ligand and electron donor in the system. Also from the Fig 2, we could prove that Fe^{II} complex catalyst is superior to Fe^{III} complex since the approach for Fe^{II} to be reduced to Fe^I is easier.

However, FeCl₃ is more stable and easier to save compare to FeCl₂, thus FeCl₃ seems to be more superior as catalyst scoure in this system. When FeBr₃ and FeSO₄ as catalyst source for photocatalytic performance, 12.4 and 10.3 TON are obtained respectively, these indicate that counter ions were shown to play an important role in governing the degree of Photocatalytic CO₂ to CO. Such dependence of Photocatalytic CO₂ to CO on the nature of the anions may be explored in the development of other Photocatalytic system.



The electronic absorption spectra at different irradiation times: [Ru](4×10⁻⁵ M) and solution is Fig. 3: TEOA/DMF = 1/5(v/v), A without and B with FeCl₃(1×10⁻⁵ M). Xe lamp through filters giving light within the scope of $410 \le \lambda \le 750$ nm.

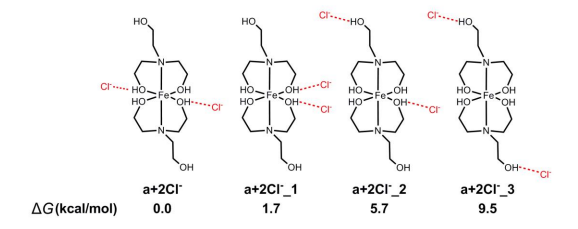


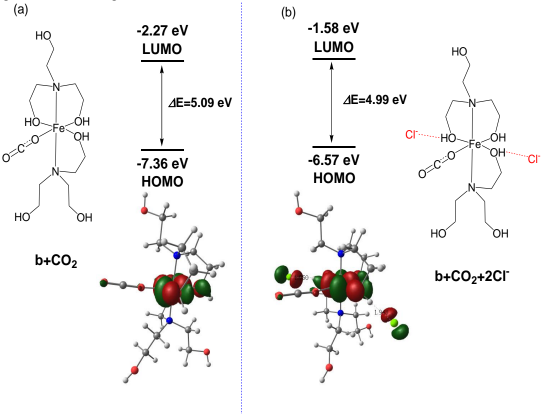
Fig 4:The relative stability of four kinds of structure for Chlorine ion couple with complexes in different positions in DMF.

As shown in Fig 5, when CO₂ was bonded to the complex's central metal, the energy band gap was 4.99 eV in the reaction if the Cl was involved. However, when the energy band gap was 5.09 eV, if the Cl⁻ was not involved. The difference between the values was 0.1 eV, which also proved that the reaction was more favorable when external ions were involved. Therefore, external ions were crucial in the photocatalytic reduction of CO₂, and could influence the overall outcome of the reaction.

Additionally, Cl⁻ in the complexes played an important role in CO₂ reduction. As shown in Fig 6, bond length stretched by 0.008~0.016 Å when different counter anions were coupled with the central metal in DMF. These results were consistent with the experimental results, where the catalytic effect of central Fe(II) metal with halogen ions, especially Cl⁻, was superior. The results showed that when SO₄²coordinated with the complexes, the bond length reached a maximum of 2.056 Å, and therefore, bond cleavage occurred allowing CO2 to attack the central

and undergo reduction. However, ion [Fe^{II}(TEOA)₂]SO₄ catalyst was not optimal for CO₂ catalytic reduction, because the volume of SO₄²⁻ in its spatial structure was much larger than that of other external ions in the experiment. This created vast steric hindrance, making it very difficult for CO₂ to attack.

On the basis of the present work and previous results [18], and according to previous literature reports [19], the process of Fe(III) to Fe(II) is a rapid process during the photocatalytic system, and the key catalytic step is the combination of Fe(I) with carbon dioxide, hence, we propose the following mechanism (Scheme 1). In the lower Fe oxidation state, one CO₂ are attached to Fe, then by a three proton and four electron (PCET) process, H₂ and CO are generated, meanwhile, the Fe catalyst are regenerated. Among these five Fe catalysts, the catalysts with chloridion as the external ions, Fe^{III}(TEOA)₂Cl₃ and Fe^{II}(TEOA)₂Cl₂ have better catalytic effect.



Iso-grams and HOMO and LUMO orbital energies of Fe(TEOA)₂ bonded to CO₂ without external Cl Fig. 5: involvement (a) and with external Cl⁻ involvement (b).

Fig. 6: The change of bond length when different counter anions couple with the central metal in DMF.

Fe(III/II) external ions
$$C\Gamma \text{ (best)}$$

$$B\Gamma \text{ Fe(II)} + CO + OH^- + H_2$$

$$2Ru(bpy)_3^{2+*} 2Ru(bpy)_3^+ Fe(I)$$

$$X \text{ Fe(II)} + CO + OH^- + H_2$$

$$XH^+$$

$$X = BIH \text{ or TEOA}$$

$$OC$$

$$OH$$

Scheme-1: Proposed mechanism for the photosensitized Catalytic reduction of CO₂ to CO.

Conclusions

In conclusion, we investigated the effect of external ions on the photocatalytic reduction of CO₂ to CO using several Fe-based homogeneous catalysts. The photocatalytic systems are irradiated under visible light (410 $\leq \lambda \leq$ 750 nm). All catalysts are easily prepared or self-assembled. During photocatalytic CO₂ reduction, the catalysts maintains activity probably due to the superfluous TEOA in the system that restrains its dissociation.

The presented photocatalytic systems are simple, where CO production over 10 h occurs with a TONs of ca. 89 for CO is obtained (negligible amount of H₂ is formed). Based on experiments and calculations, Different external counterions were proved to have different effects in governing the degree of photocatalytic CO2 to CO. These results are significant for the preparation of photocatalysts, provided reference for the selection of external ions.

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