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## Highly efficient CO<sub>2</sub> reduction under visible-light on noncovalent Ru—Re assembled photocatalyst: evidence on the electron transfer mechanism

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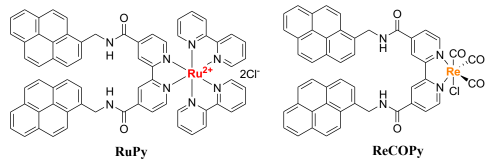
Keywords: Artificial photosynthesis, CO2 photoreduction, supramolecular assembly, electron transfer

**ABSTRACT**: Herein, two complexes: Pyrene modified photosensitizer (**RuPy**) and catalyst (**ReCOPy**) were synthetized and combined to form a novel bimetallic supramolecular assembly **RuPy**...**ReCOPy** via a combination of  $\pi$ - $\pi$  stacking and hydrogen bond interactions. This supramolecular assembly was further characterized by electrochemical and spectroscopic techniques in order to determine its redox and photochemical properties. This novel **RuPy**...**ReCOPy** supramolecular assembly was demonstrated to display a high level of performance for the photocatalytic conversion of CO<sub>2</sub> conversion into CO with turn over frequency and CO quantum yield much higher (TOF: 18 h<sup>-1</sup>, Φ<sub>CO</sub>: 0.77) than the value obtained for the benchmark system Ru(bpy)<sub>3</sub>Cl<sub>2</sub> (**Ru**) with Re(bpy)(CO)<sub>3</sub>Cl (**ReCO**) (5.3 h<sup>-1</sup>, Φ<sub>CO</sub>: 0.38). Time resolved spectroscopy and Density Functional Theory calculations revealed that an electron transfer mechanism is at the origin of the highly photocatalytic performance of **RuPy**···**ReCOPy**.

#### 1. INTRODUCTION

In the recent years, the use of CO<sub>2</sub> as an alternative carbon source has become increasingly attractive for the production of useful carbon compounds, i.e. HCOOH, MeOH, CH<sub>4</sub> and CO, among others. [1-3] In particular, CO<sub>2</sub> reduction using photocatalysts to mimic the solar light driven photosynthesis is a topical subject. In this field, the photocatalytic systems envisaged so far are split into three categories: *i*-semiconductor systems, [4-8] *ii*- biohybrid systems [9-12] and *iii*- organometallic complexes. This later strategy was initiated by Lehn et *al*. in 1983 using Re(bpy)(CO)<sub>3</sub>Cl catalyst to convert CO<sub>2</sub> into CO. [13,14] A wide range of metal-based complexes, i.e. Ru, Rh, Co and Mn have been further demonstrated to show even higher photocatalytic activities for CO<sub>2</sub> conversion.[15,16] Generally, a photosensitizer (PS) is required in tandem with the co-catalyst (Cat) to enable absorption of visible light and transfer the electron from a sacrificial electron donor to the Cat. Typically, Ishitani *et al*. reported several supramolecular photocatalysts by covalent linking of different photosensitizers and co-catalysts. In these systems, an efficient and faster electron transfer (ET) via intramolecular process was demonstrated to impact substantially their catalytic performance. [17-25] A faster ET from the photosensitizer to the metallic center

of the catalyst reduces the undesirable deactivation process of the PS excited state. Reducing the distance between the PS and the Cat via non-covalent interactions is an alternative promising strategy developed recently for making the ET more efficient. This allows adjusting the PS/Cat ratio, simplifying the synthesis process comparing to covalently linked PS-Cat, expanding their use and adaptation with other PS/Cat moiety and/or their potential adsorption on conductive support, etc. In this context, Kubiak et al. reported, recently, intramolecular ET by assembling Ru(dac)(bpy)<sub>2</sub>(PF6)<sub>2</sub> and Re(dac)(CO)<sub>3</sub>Cl (dac = 4,4'-bis(methyl acetamidomethyl)-2,2'-bipyridine, bpy = 2,2'-bipyridine) via (-NH---O=C-) hydrogen bond interactions resulting in a substantial improvement of the CO<sub>2</sub> conversion as compared to the combination of Re(dac)(CO)<sub>3</sub>Cl with the commercial Ru(bpy)<sub>3</sub>PF<sub>6</sub>.[26,27] Another system stabilized by  $\pi$ -  $\pi$  stacking interactions was reported, recently, by He et al, with Re[4,4'-di(pyren-1-yl)-2,2'-bipyridine](CO)<sub>3</sub>Cl (4.4'dipyrenyl-Re) as co-catalyst in the presence of Ru(bpy)<sub>3</sub>Cl<sub>2</sub> enabling an efficient conversion of CO<sub>2</sub> into CO.[28] This intermolecular  $\pi$ - $\pi$  stacking was found to boost ET leading to a turnover number (TON) twice higher than that obtained using the commercial Re(bpy)CO<sub>3</sub>Cl. On the other hand, the 4.4'-dipyrenyl-Re was active for CO<sub>2</sub> photoreduction without the photosensitizer under solar light. This led to a light harvesting enhancement due to the  $\pi$ -  $\pi$  conjugation between the pyrene and the bipyridine groups of the catalyst as well as to the accelerated intermolecular ET through the stacked pyrenes. Very recently, Lu et al. reported a series of Ru(Phen)<sub>3</sub><sup>2+</sup> complexes (Phen: 1, 10-Phenanthroline) modified by pyrene as photosensitizers in association with cobalt-complex based catalyst for CO<sub>2</sub> photoreduction. [29] The significant enhancement, observed with one of the modified photosensitizer, was assigned to an improvement of their photosensitization due to a long-lived excited state and suitable excited-state oxidation potential. It is also well documented that the  $\pi$ - $\pi$  interactions enable the immobilization of pyrene modified catalysts on the surfaces of carbon allotropes for recycling purposes, i.e. reduced graphene oxide, carbon nanotubes and C60 among others. [30-34] Typically, Gray et al. reported the synthesis of a series of bipyridine-based complexes containing two pyrenes moieties to be immobilized on carbon electrode via  $\pi$ - $\pi$  stacking for electrochemical CO<sub>2</sub> reduction. [35,36] Herein, we report two ruthenium and rhenium-based complexes, i.e. RuPy and **ReCOPy** respectively, containing both, two pyrenes linked via -CH<sub>2</sub>-NH-CO- bonds with the bipyridine moieties (Fig. 1). The so-designed complexes are assembled to produce a photocatalytic supramolecular PS/Cat system via an unconventional combination of  $\pi$ - $\pi$  stacking and hydrogen bond interactions. In addition to the short separating distance between PS and Cat, we demonstrated that the  $\pi$ - $\pi$  stacking promotes an efficient ET mediated by the pyrene moieties to achieve an outstanding conversion of CO<sub>2</sub> under visible light. Time resolved photoluminescence experiments were further combined with Density Functional Theory (DFT simulations) to shed light into the efficient electron transfer at the origin of the remarkable performances of this novel supramolecular photocatalytic system.



**Fig. 1. ReCOPy** and **RuPy** complexes combined in this work to design a novel supramolecular assembly via hydrogen bond and  $\pi$ - $\pi$  stacking.

#### 2. EXPERIMENTAL SECTION

#### 2.1. Materials

1,8-diazabicyclo[5.4.0]undec-7-ene (DBU), 4,4'-Dimethyl-2,2-bipyridine and the complex Re(bpy)(CO)<sub>3</sub>Cl were purchased from TCI chemicals. Ru(bpy)<sub>2</sub>Cl<sub>2</sub>.6H<sub>2</sub>O, Ru(bpy)<sub>3</sub>Cl<sub>2</sub>, ReCO<sub>5</sub>Cl, DMF, SOCl<sub>2</sub> and 1-pyrenemethylamine hydrochloride were purchased from Merck company. Commercially available acetonitrile, toluene, THF, DMF were used without any distillation.

#### 2.2. Complexes RuPy and ReCOPy preparation

#### 2.2.1. 4,4'-Dicarboxylic acid-2,2-bipyridine (HOOC-Bpy-COOH).

4,4'-Dimethyl-2,2-bipyridine (5.0 g) in heated  $H_2SO_4$  (125 mL, 70-80 °C) was oxidized by adding solid potassium dichromate (24 g) slowly. The temperature was held consistently between 70 and 80 °C during the transfer. The deep green mixture was then poured over 800 mL of ice/  $H_2O$ , affording a light-yellow precipitate that was isolated by vacuum filtration and was washed with  $H_2O$ . The solid was then refluxed in 50%  $HNO_3$  (150 mL) for 4 h. The cooled solution was then poured over ice and diluted with 800 mL of  $H_2O$ . A white powder was isolated by vacuum filtration and washed with  $H_2O$  (5 g, 73%). <sup>1</sup>H NMR (400 MHz, DMSO- $H_2O$ ) 8 (ppm): 13.83 (2H, br s, R-COOH), 8.92 (d, J=5.1 Hz, 2H), 8.85 (s, 2H), 7.92 (dd, J=4.9, 1.2 Hz, 2H).

### $2.2.2. \quad N^{4,4'} - bis(1-pyrenylmethyl) - [2,2'-dipyridyl] - 4,4'-dicarboxamide (Ligand-Bpy-Py). \\$

The acid chloride of 2,2′- bipyridine-4,4′-dicarboxylic acid was prepared by refluxing 0.37 g (1.5 mmol) of the 4,4-dicarboxylic acid-2,2-bipyridine acid in 15 mL of thionyl chloride for 6 h. Remaining thionyl chloride was removed under vacuum. In a separate flask, 0.8 g (3 mmol) of 1-pyrenemethylamine hydrochloride were combined with 150 mL of dry acetonitrile and 0.9 mL of 1,8-diazabicyclo[5.4.0]undec7-ene (DBU). The resulting solution was transferred to the solid bipyridine-acid chloride via cannula. The mixture was stirred for 12 h at room temperature, at which time 20 mL of saturated sodium bicarbonate and

130 mL of distilled water were added to form a beige precipitate. The precipitate was collected by filtration and washed with deionized water. The precipitate was dried under vacuum to give 0.62 g (61%) of an off-white solid.  $^{1}$ H NMR (400 MHz, DMSO-d6)  $\delta$  (ppm), 9.73 (t, J = 3.6 Hz, 2H), 8.86-8.84 (m, 4H), 8.51, (d, J= 6Hz, 2H), 8.34-8.26 (m, 9H), 8.18 (m, 5H), 8.13 (d, J= 5.3 Hz, 2H), 8.08 (t, J = 4.4 Hz, 2NHamide), 5.28 (d, J= 3.7 Hz, 4H).

## 2.2.3. N<sup>4,4'</sup>-bis(1-pyrenylmethyl)-[2,2'-dipyridyl]-4,4'-dicarboxamide)-bis(2,2'-bipyridine dichloro ruthenium (II) (RuPy).

A solution of Ru(2,2'-bipyridine)<sub>2</sub>Cl<sub>2</sub> (21 mg, 0.044 mmol) with **Ligand-Bpy-Py** (30 mg, 0.044 mmol) in 0.6 mL of DMF was refluxed 18 h at 153 °C under argon. The product was precipitated by addition of diethyl ether and sonicated during five minutes. The product was collected by centrifugation and washed with additional diethyl ether. The obtained solution was then dried by rotary evaporation to obtain 26 mg of dark red powder. (Yield = 50%) <sup>1</sup>H NMR (400 MHz, DMSO-d6) δ (ppm) 9.95 (t, J = 3.5 Hz, 2H), 9.40 (s, 2H), 8.83 (t, J = 5.2 Hz, 4H), 8.52 (d, J = 6.4 Hz, 2H), 8.33-8.23 (m, 9H), 8.20-8.12 (m, 9H), 8.09 (t, J = 5 Hz, 2H), 7.96-7.89 (m, 4H), 7.74 (d, J = 3.6 Hz, 2H), 7.68 (d, J = 3.6 Hz, 2H), 7.54-7.51 (m, 2H), 7.46-7.43 (m, 2H), 5.30 (d, J = 4.4 Hz, 4H). <sup>1</sup>H NMR (125 MHz, DMSO-d6) δ (ppm) 163.13 (CO), 157.67 (C<sub>IV</sub>), 156.81(C<sub>IV</sub>), 156.76 (C<sub>IV</sub>), 152.52 (CH), 152.09 (CH), 151.53 (CH), 141.73 (C<sub>IV</sub>), 138.67 (CH), 138.63 (C<sub>IV</sub>), 132.57 (C<sub>IV</sub>), 131.21 (C<sub>IV</sub>), 130.71(C<sub>IV</sub>), 130.69(C<sub>IV</sub>), 128.60 (C<sub>IV</sub>) 128.44 (CH), 128.41 (CH), 128.14 (CH), 127.83 (CH), 127.67 (CH), 126.77 (CH), 126.25 (C<sub>IV</sub>), 125.81 (CH), 125.66 (CH), 125.21 (CH), 125.01 (CH), 124.48 (C<sub>VI</sub>), 124.33 (C<sub>IV</sub>), 123.87 (CH), 122.55 (CH), 41.74 (CH<sub>2</sub>). ESI-MS (positive) m/z: found 542.14 (**RuPy**/2 – 2Cl), calculated 539.14 (**RuC**<sub>66</sub>**H**<sub>46</sub>**N**<sub>8</sub>**O**<sub>2</sub>)<sup>2+</sup>.

# 2.2.4. fac-(N<sup>4,4'</sup>-bis(1-pyrenylmethyl)-[2,2'-dipyridyl]-4,4'-dicarboxamide)tricarbonylcholoro rhenium (I) (ReCOPy).

A solution of Re(CO)<sub>5</sub>Cl (18 mg, 0.049 mmol) with **Ligand-Bpy-Py** (30 mg, 0.044 mmol) in 1 mL of toluene was refluxed 18 h at 110°C under argon. The solution was centrifuged to give an orange solid. The solid material was redissolved in THF before being centrifuged once more. The obtained solution was then concentrated and dried by rotary evaporation to obtain 30 mg of an orange precipitate. (Yield = 68%). <sup>1</sup>H NMR (400 MHz, DMSO-d6) δ (ppm) 9.92 (t, J = 3.7 Hz, 2H), 9.18 (d, J= 3.8 Hz, 2H), 9.12 (s, 2H), 8.46 (d, J= 6.1 Hz, 2H), 8.34-8.24 (m, 9H), 8.07-8.18 (m, J= 5.27Hz, 9H), 8.08 (t, J= 5 Hz, 2NH amide), 5.32 (d, J= 4 Hz, 4H). ESI-MS (positive) m/z: found 999.14 (**ReCOPyNa**), calculated 999.13 (**ReC**<sub>49</sub>**H**<sub>30</sub>**N**<sub>4</sub>**O**<sub>5</sub>**ClNa**).

#### 2.3. Electrochemistry

Electrochemical measurements were realized with a PowerEase 300W Power Supply, 230 VAC. All experiments were performed in DMF, the supporting electrolyte was 0.1 M tetrabutylammonium

hexafluorophosphate (TBAPF<sub>6</sub>), using a standard three-electrode configuration: 2 mm diameter. Platinum disk working electrode, a silver wire immersed in a solution of 0.1 M TBAPF<sub>6</sub> served as a reference electrode and the counter electrode was a Pt wire. The ferrocenium/ferrocene ( $Fc^{+/0}$ ) couple was used as an external reference in all experiments to calibrate potentials. Voltammetry collected for solution-soluble redox couples with typically concentrations of  $3 \times 10^{-4}$  M with scan rate of 100 mV/s.

#### 2.4. Photocatalytic tests

The photocatalytic  $CO_2$  reduction was tested in an in-situ reactor developed in the LCS laboratory and presented in Fig. S5. The reactor is made from glass with horizontal tubes which ensuring the FTIR beam pathway through  $CaF_2$  windows. The reactor is equipped with air circulator system for cooling and two gas inlet and outlet. The visible irradiation is ensured from the top through a removable quartz window. The quartz and  $CaF_2$  windows are attached to the reactor with screw fittings and the tightness is obtained by adapted Teflon O-rings. The total internal volume of the reactor is  $160 \pm 5$  ml. All IR spectrum measurements with the in situ FTIR reactor were monitored in real time with 5 min per spectrum of time resolution. IR spectra of the reactor headspace were recorded using Nicolet 6700 IR spectrometer (Thermo Fisher 14 Scientific) equipped with an MCT detector. All photocatalytic tests presented in this work were performed in batch configuration and at room temperature. As a light source, Xe-lamp 16 (LC8 Hamamatsu, 200 W) with pass-high filter at 390 nm was used. Complexes were dissolved in DMF/H<sub>2</sub>O mixture (10/1.8 ml) and the solution was bubbled with saturated with  $CO_2$  for one hour with a flow rate of 8 ml min<sup>-1</sup>. Then, triethanolamine (TEOA) was introduced as electron donor (220 mg in 1 mL) in the reactor where the  $CO_2$  bubbling was maintained for additional 15 minutes.

#### 3. RESULTS AND DISCUSSION

#### 3.1. Electro-photochemical characterizations

Two complexes were synthesized in this study, namely **ReCOPy** and **RuPy** (**Fig. 1c**). The first one was prepared according to the protocol reported in a previous study which considered this complex for the electro-catalytic reduction of CO<sub>2</sub>. [36] **RuPy** was synthetized, for the first time in this work, from the equimolar mixture of Ru(bpy)<sub>2</sub>Cl<sub>2</sub> and **Bpy-Py** ligand. The other two commercial homologues **Ru** and **ReCO** complexes were used for comparison.

The electrochemical analysis was performed in order to determine the redox potentials of the different systems. The cyclic voltammetry (CVs) of **Ru** and **ReCO** complexes agrees well with the literature. [27] A first reduction from Ru<sup>2+</sup> to Ru<sup>+</sup> and from Re<sup>+</sup> to Re<sup>0</sup> occurs at -1.77 V and at -1.79 V, respectively, with reversible oxidation bands at -1.71 V and -1.70 V, respectively (**Fig. 2**). Regarding the **ReCOPy** complex, the reduction band of **ReCO**<sup>+</sup>/**ReCO**<sup>0</sup> is shifted to -1.96 V with a quasi-reversible oxidation band at -1.86V.

On the other hand, The CV analysis of the **RuPy** complex showed a first oxidation band of **RuPy**<sup>2+</sup>/**RuPy**<sup>+</sup> at -1.73V. In addition to the UV-visible absorption data (bad gap values), these reduction potential values will be used later for calculating the energetic positions of the HOMO (Highest Occupied Molecular Orbital) and LUMO (Lowest Unoccupied Molecular Orbital) of the corresponding complexes.

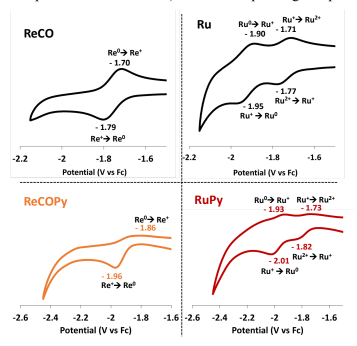


Fig. 2. Cyclic voltammetry for the ReCO, ReCOPy, Ru and RuPy complexes in DMF. Conditions: 100 mV/s, 0.1 M TBAPF<sub>6</sub>, Pt working electrode, Pt wire counter electrode, Ag/AgCl reference electrode. Referred to ferrocenium/ferrocene couple.

The UV-Vis spectra of the complexes highlight the interaction of pyrene function with the metal center (**Fig. 3a**). The absorption bands, located at 458 nm and 370 nm for **Ru** and **ReCO**, respectively, are assigned to the metal to ligand charge transfer (MLCT). [37,38] However, a red-shift of ca. 20 nm is noted in MLCT absorbance range for both **RuPy** and **ReCOPy** complexes (478 and 405 nm, respectively). This red-shift consolidates the hypothesis of the electron delocalization between the metal center and the pyrene indicating strong interactions between them. It should be noted that no significant influence of the grafted pyrene group on the absorbance coefficient was observed in this work, in contrast to what has been observed when pyrene is directly linked to the bipyridine group of Re(bpy)(CO)<sub>3</sub>Cl. [28] In the last case, a significant increase in the coefficient molar absorption of the MLCT is observed in addition to the red-shift. This difference can be explained by a higher delocalization of the electron between the directly-linked pyrene and the 4,4'-dipyrenyl-Regroup via a single bond. In contrast, the contribution of the pyrene in the electron delocalization is probably lower for **ReCOPy**, however the structure flexibility is higher owing to the presence of -CH<sub>2</sub>-NH-CO- linker, which is more suitable, in our case, for promoting the desired interaction with **RuPy** photosensitizer. In order the investigate the interaction between **RuPy** and **ReCOPy**, a mixture of ratio 1:1

of complexes was analyzed by UV-Vis ( $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}_{exp}$ ) and the obtained spectra was compared with that of theoretical calculated from the addition of the individual  $\mathbf{RuPy}$  and  $\mathbf{ReCOPy}$  spectra ( $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}_{ealc}$ ). (**Fig. 3b**) The subtraction of the two spectra  $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}_{exp}$ - $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}_{ealc}$  indicates the presence of band-edge shifts with a modification in some band intensity. 321 and 339 nm resulting from the band-shift of pyrene at 330 and 345 nm, respectively. On the other hand, the characteristic bands 278 and 345 nm of pyrene was more intense in the  $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}_{exp}$  spectrum. In addition, a bleaching were observed at 473 and 350 nm and attributed to the MLCT and metal centered d-d transitions (MC), respectively. This bleaching is accompanied with a relative increase of the band intensity centered at 290 nm and assigned to  $\pi$ - $\pi$ \* transitions (\*LC) of the ligand. [39] These observations confirm the interaction between  $\mathbf{RuPy}$  and  $\mathbf{ReCOPy}$  in solution through the pyrene moieties which indicate a plausible formation of supramolecular assembly of  $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}$ . It should be noted, that the relative low intensity due to the interaction could probably due to the fact of the  $\mathbf{ReCOPy}$ ... $\mathbf{ReCOPy}$  interaction. Therefore, the difference, is the additional impact of the  $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}$  interaction in respect to the self- $\mathbf{ReCOPy}$  interaction.

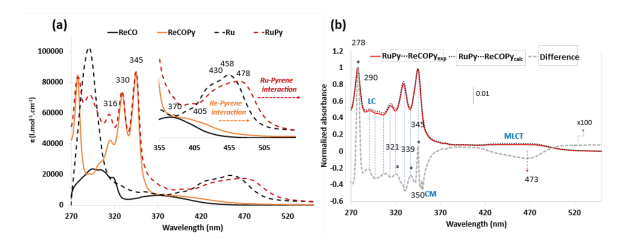


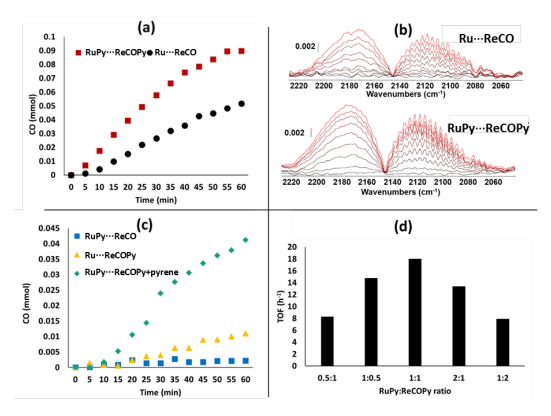
Fig. 3. (a) UV-Visible spectra of Ru, RuPy photosensitizers and of ReCO, ReCOPy co-catalysts in DMF. Insert: zoom on the region of the MLCT bands. (b) Normalized spectra of RuPy-ReCOPy (1:1) mixture (RuPy-ReCOPy<sub>exp</sub>) and of mathematic addition of RuPy and ReCOPy (RuPy···ReCOPy<sub>calc</sub>) with the difference between the two spectra, respectively. [Ru] = [RuPy] = [ReCO] = [ReCOPy] = 0.4 mM, [RuPy···ReCOPy<sub>exp</sub>] : [RuPy] = [ReCOPy] = 0.2 mM \*characteristic bands of pyrene

#### 3.2. Photocatalytic performance.

All photocatalytic tests were carried out under similar visible light irradiation condition, in DMF, using triethanolamine (TEOA) as sacrificial electron donor and after saturation of the catalytic batch reactor with CO<sub>2</sub>. The gas phase was monitored during the reaction by *in-situ* FTIR and the amount of CO produced was quantified following the characteristic bands at 2116 and 2171 cm<sup>-1</sup> (**Fig. 4b**) as reported in our recent works. [40] It should be noted that the CO is formed selectively from the CO<sub>2</sub> reduction as demonstrate Figure S8

using labelled CO<sub>2</sub> ( $^{13}$ CO<sub>2</sub>). All tests were reproduced at least two times and the error bars were estimated to be less than  $\pm 10\%$  (**Fig. S9**). The reactor temperature was maintained at around 25°C by air cooling circuit. The selectivity toward CO vs HCOOH formation was >99% in all cases, under our reaction conditions as further confirmed by gas chromatography (**Fig. S9**).

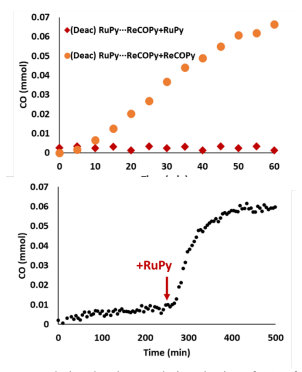
An equimolar mixture of complexes RuPy...ReCOPy (1:1) was found to produce around 0.1 mmol of CO after one hour with an initial reaction rate  $V_0$  of 0.12 mmol.h<sup>-1</sup> and turn over frequency (TOF) of  $18\pm2$  h<sup>-1</sup>. These results were compared to those obtained with the commercial Ru and ReCO complexes, tested under the same conditions. Remarkably, our new **RuPy**...**ReCOPy** s system shows a substantially higher activity with a TOF three times higher than the value obtained for Ru...ReCO (18 vs 5.3 h<sup>-1</sup>, respectively) as well as four-fold higher initial reaction rate  $V_0$  (0.12 vs 0.03 mmol.h<sup>-1</sup>, respectively). (**Table S1**) Additionally, a very high quantum yield ( $\Phi_{co}$ ) for CO formation was calculated to be 0.77 for **RuPy...ReCOPy** and 0.38 in case of Ru-ReCO .Comparing the relative TOF and Φ<sub>co</sub> with that reported in the literature when mononuclear homogeneous counterparts (PS + Cat) are used as reference allows ranking our system as one of the four most efficient counterpart systems reported in the literature (Table S2). This observation emphasizes a major role of the pyrene function on TOF and reaction rate in the photocatalytic process (Fig. 4a). However, the absorbance behaviors of the photosensitizers as well as the redox potentials of the different complexes cannot justify this gap between both systems. In order to better understand this performance, equimolar mixtures of PS (Ru or RuPy) and Cat (ReCO or ReCOPy) binary combinations were further explored and tested under similar conditions. The combination of RuPy...ReCO was tested with 1:1 ratio and surprising negligible activity was detected (TOF =  $0.4 \, h^{-1}$ ). Moreover, **Ru**...**ReCOPv** (1:1) showed a poor conversion (TOF =  $2.2 \text{ h}^{-1}$ ) (Fig. 4c). These results seem contradictory to what has been reported recently on the benefit effect of the pyrene-modified photosensitizer [28,29] where a higher efficiency of Ru...ReCOPy systems (as well as for RuPy...ReCO) is expected with respect to Ru...ReCO. The very low activity indicates that ET between the pyrene modified RuPy/ReCOPy structures and unmodified Ru/ReCO structures are not favorable despite the suitable redox potentials of the different couples. The effect of PS:Cat ratio on the photocatalytic activity of the system was also investigated. The tests with different RuPy:ReCOPy ratios were performed under similar conditions and the results are reported in Fig. 4d and S12. A lower concentration of the PS or Cat (ratio 0.5:1 and 1:0.5) led to a significant decrease of the TOF (8.3 h<sup>-1</sup> and 14.8 h<sup>-1</sup> respectively). Moreover, a higher concentration of the photosensitizer (RuPy:ReCOPy 2:1) or of the catalyst (RuPy:ReCOPy 1:2) causes a dramatic drop of the performance (TOF =  $13.4 \text{ h}^{-1}$  and  $7.9 \text{ h}^{-1}$ , respectively). These observations are different from that obtained with a covalent supramolecular PS-Cat where the highest activities were observed with 1:3 and 1:2 of PS:Cat ratios. [41-43] In order to figure out the role of the linked pyrene function on the complexes, a complementary experiment was performed by adding free pyrene (4 equivalent relative to the PS) to the **RuPy**...**ReCOPy** with the idea that free pyrene can interact with the pyrenes of the complexes and consequently minimize the possible  $\pi$ - $\pi$  interactions between the **RuPy** and **ReCOPy** (It should be noted that free pyrene couldn't interfere with the light absorption in the visible irradiation range). As excepted, a decrease of the catalytic activity was observed (**Fig. 4c**), with two times lower TOF (8.2 h<sup>-1</sup>) than that observed for pyrene-free system and close to that observed for **Ru**...**ReCO** complex used as reference.



**Fig. 4.** Evolution of the CO amount during the photocatalytic reduction of CO<sub>2</sub> under visible light over: a) **RuPy**...**ReCOPy** and **Ru**...**ReCO**; c) **Ru**...**ReCOPy**, **RuPy**...**ReCO** and **RuPy**...**ReCOPy**+pyrene; d) Evolution of the TOF vs the ratio of **RuPy**:**ReCOPy**. b) Characteristic IR CO band evolutions in the first 30 minutes of irradiation for **RuPy**...**ReCOPy** and **Ru**...**ReCO**. Reaction condition: [**Ru**]=[**RuPy**]= 0.38 mM; [**ReCO**]=[**ReCOPy**)] = 0.38 mM; [TEOA]= 0.12 M; [H<sub>2</sub>O] = 8.3 M; Vol<sub>DMF</sub>= 11 ml; Xe-Lamp; Irradiance~10 mW/cm²; Filter pass high for  $\lambda$ >390 nm; errors estimated lower than 10% on three reproducible experiments. TOFs were always based on the amount of Re complexes.

It to be noted that our new system RuPy...ReCOPy (1:1) displays a lower stability with respect to Ru...ReCO system in these conditions. A total deactivation was observed after ≈70 minutes (Fig. S10). In order to clarify the origin of this deactivation, a set of additional experiments was performed, e.g. the CO₂ purging and/or introduction of RuPy or ReCOPy moiety after the total deactivation. We evidenced that the activity can be only recovered by adding a fresh Cat (ReCOPy) to the solution showing that the degradation or the poisoning of this Cat is the only responsible of the deactivation process (Fig. 5a). However, the synergistic increase of the activity after adding "fresh" RuPy to solution containing only ReCOPy, and

previously irradiated for 240 minutes under CO<sub>2</sub> atmosphere in presence of TEOA, discarded the hypothesis of the catalyst photo-degradation (**Fig. 5b**). Therefore, the poisoning of this latter seems to be more realistic and probably due to the formation of a sub-product obtained from a possible reaction of the catalyst with the sub-product of electron donor (e.g. TEOA'). [44]. These experiments highlighted further the nature of the **RuPy/ReCOPy** interactions. The synergistic activity observed after adding the fresh **ReCOPy** to **RuPy**...**ReCOPy** or **RuPy** to the **ReCOPy** revealed the reversibility and the advantage of the non-covalent nature of the interactions between the PS and the Cat.

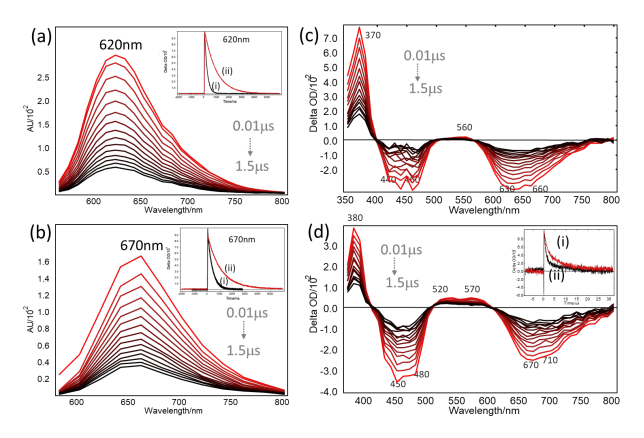


**Fig. 5**. a) Evolution of the CO amount during the photocatalytic reduction of CO<sub>2</sub> after 2 hours of bubbling CO<sub>2</sub> on deactivated **RuPy**...**ReCOPy** system and adding of **RuPy** and **ReCOPy**. (b) Evolution of the CO amount during the photocatalytic reduction of CO<sub>2</sub> over **ReCOPy** system before and after adding (indicated with arrow) of **RuPy** photosensitizer.

#### 3.3. Time resolved emission of <sup>3</sup>MLCT and <sup>3</sup>IL excited state of PS.

Adjusting the excited-state lifetime of the photosensitizers is a key factor for an efficient reduction of CO<sub>2</sub> over PS/Cat system. Hence, the excited-state properties of **Ru** and **RuPy** were explored by measuring the time resolved photoluminescence (TR-PL) and absorption transient (AT) (**Fig. 6**). The PL of **Ru** and **RuPy** were significantly quenched as exposed to air (**Fig. 6a**), indicating the phosphorescence emission process, which mainly derived from the triplet state. <sup>45</sup> The TR-PL spectrum of **Ru** showed a typical <sup>3</sup>MLCT emission at 620 nm in DMF (**Fig. 6a**; **Fig. S13**). The <sup>3</sup>MLCT emission of **RuPy** is centered at 670 nm (**Fig. 6b**, **Fig. S13**), red-shifted for 50 nm with respect to **Ru**. This red-shift confirms the strong electrostatic interactions

between the Ru-center and pyrenyl, in agreement with the UV-visible absorbance results. These results are confirmed by the absorption transient. For the **Ru** (Fig. 6c), absorption transient (AT) bands centred at 370 nm, and 560 nm and assigned to  $T_1$  and  $T_2$  transients, respectively are observed, with  $\lambda_{\rm exc}$ =532nm (as well as with  $\lambda_{\rm exc}$ =355nm), with a similar kinetic decay of the bleaching (at 450-460 nm, 630 and 660 nm). For RuPy, the corresponding AT bands are red-shifted, in agreement with the PL and UV-visible absorbance, and centered at 380 nm and 570 nm with a similar kinetic decay of the bleaching (at 450-480 nm, 670 and 710 nm). The bleaching in the 600-700 nm region is mainly assign to the phosphorescence emission due to radiative relaxations of the triplet state. The lifetimes of the PL, AT and bleaching are very close for Ru as well as for RuPy. The kinetic decay trace of the RuPy's PL and AT revealed 1.3-1.5 times higher excitedstate lifetime than that of <sup>3</sup>Ru for a similar concentration (1.08-1.30 µs vs 0.75-0.90 µs, respectively). The <sup>3</sup>RuPy was also demonstrated to be 2.5 times less sensitive toward O<sub>2</sub> (Table 1). More surprising, a new distinctive AT band at 520 nm was selectively observed for RuPy with a longer lifetime of 35 µs (Insert of Fig. 6d). The sensitivity of this AT toward oxygen confirms that it corresponds to a triplet state and can probably be correlated to a formation of an exciplex <sup>3</sup>(RuPy...RuPy) and/or to an intermolecular charge separation. Moreover, the quenching of <sup>3</sup>RuPv by TEOA was found 2.8 times higher than that for <sup>3</sup>Ru (2.11×10<sup>6</sup> M<sup>-1</sup>s<sup>-1</sup> vs 7.4×10<sup>5</sup> M<sup>-1</sup>s<sup>-1</sup>, respectively), indicating a faster electron transfer from the electron donor to the RuPy. It should be noted that the quenching rate constants calculated from the emission (Fig S11) are very similar to that calculated from the AT. A characteristic shoulder with a very short lifetimes (<10 ns) was selectively observed for RuPy (Fig. S13a), representative of a dual emission emerged in the PL spectrum. The PL emission spectra of RuPy after excitation at 355 nm confirmed this inspection (Fig. S13bd) with additional emissions centred at 400 nm and 520 nm. This emission could be assigned to the pyrene moiety. Consequently, PL studies of pyrene molecule and of the RuPy's ligand were performed. The PL measurement results indicated that this emission corresponds to the intra-ligand triplet excited state (<sup>3</sup>IL). Therefore, observing this <sup>3</sup>IL upon 532 nm excitation (no absorption of pyrene and the ligand at 532 nm) substantiates the electron/energy transfer between the Ru-center to the pyrene part. The significant decrease of the <sup>3</sup>IL lifetime of pyrene (Py), in the following order Py(200ns)<ReCOPY(120 ns)<Bpy-Py(100 ns)<RuPy(85ns), confirms this assumption (Fig. S13g).



**Fig. 6.** (a) and (b) correspond to the nanosecond emission spectra (140ns/spectrum) of the  ${}^{3}$ MLCT of **Ru** and **RuPy**, respectively. In insert: normalized kinetic decay trace of  ${}^{3}$ MLCT emission of **Ru** and **RuPy** monitored at 620 nm and 670 nm, respectively, under air (i) and argon (ii) atmospheres. **(c)** and **(d)** report the evolution vs the time (140ns/spectrum) of the absorption transition (AT) of **Ru** and **RuPy**. Insert of Fig. d: normalized kinetic decay traces of the AT at 520 nm (i) and 570 nm (ii).  $\lambda_{\rm exc} = 532$  nm, [**RuPy**]=1.7 mM; [**Ru**]= 3 mM; all spectra were recorded under Ar at 298K.

We should keep in mind that the electron transfer is promoted only after the reduction of the  ${}^{3}\text{PS}$  to PS which is achieved by an electron transfer from the electron donor to the  ${}^{3}\text{PS}$ . Therefore, other quenching process can also play a role in the decrease of this transfer as  ${}^{3}\text{PS}$ - ${}^{3}\text{PS}$  self-quenching and  ${}^{3}\text{PS}$ -Cat quenching. The different rate constants of the  ${}^{3}\text{MLCT}$  quenching of the **Ru** and **RuPy** are reported in **Fig. S14** and summarized in **Table 1**. Interestingly, the self-quenching of  ${}^{3}\text{RuPy}$  is three times lower than that of  ${}^{3}\text{Ru}$ . However, the quenching of  ${}^{3}\text{MLCT}$  of **Ru** by **ReCOPy** is 500 times higher with respect to  ${}^{3}\text{Ru}/\text{ReCO}$ . This quenching is in part promoted by the pyrene-moiety as demonstrated from the rate constant of the  ${}^{3}\text{Ru}$  with pyrene (Table 1). The  ${}^{3}\text{Ru}/\text{Py}$  quenching is 6 times higher than that for  ${}^{3}\text{RuPy}/\text{Py}$  (and much higher than that the ET from TEOA), which could explain the very low activity of **Ru/ReCOPy** dual system. However, despite the low quenching rate of the  ${}^{3}\text{MLCT}$  of **RuPy** by pyrene, the lifetime of  ${}^{3}\text{Py}$  is dramatically affected with similar quenching rate constant founded with  ${}^{3}\text{Ru}$ . This outweighs on a  $\pi$ - $\pi$  interaction between the pyrene-moiety of **RuPy** with pyrene. On the other hand, the quenching of the  ${}^{3}\text{MLCT}$  of **RuPy** by **ReCO** is four times lower than that with **ReCOPy** which could not explain the very low efficiency of **RuPy/ReCO** 

binary system compared to **RuPy/ReCOPy**. The different data allow building the energetic diagram of the HOMO and LUMO, for the different complexes (see ESI for more details) as well as the energetic levels of the different excited states (basing on the AT experiments) of the **Ru** and **RuPy** photosensitizers (**Fig. S15**). It should be noted that the electron transfer occurs from the reduced photosensitizer \*(PS<sup>-</sup>), to the catalyst. The \*(PS<sup>-</sup>) is formed from the oxidation of the electron donor and the <sup>3</sup>PS. Therefore, a longer lifetime of **RuPy** excited state(s) is highly desired as well as a lower quenching is critical for promoting the ET from the TEOA to the <sup>3</sup>PS as well as from the reduced <sup>3</sup>PS<sup>-</sup> to the Cat. Consequently, the higher lifetime of the **RuPy** excited-state and its lower quenching with respect to that of **Ru** can justify, in part, the highest activity of the **RuPy**...**ReCOPy** system (in comparison to **Ru**...**ReCO**). However, this cannot explain the gap in the performance between **RuPy**...**ReCOPy** and **RuPy**...**ReCO**, where the energetic diagram reveals similar LUMO levels of **ReCO** with respect to **ReCOPy**. Therefore, other factors should be taken into account to elucidate this phenomenon.

**Table 1.** Stern–Volmer-quenching constants with **Ru** and **RuPy** measured at 298K from the PL kinetic decays at 620 nm and 650 nm, respectively, in presence of different quencher.

PS	${}^{3}Ru^{(a)}$	${}^{3}RuPy^{(b)}$
Q	$k_q \times 10^8 (M^{-1} s^{-1})$	$k_q \times 10^8 (M^{-1} s^{-1})$
$O_2$	299	117
Ru/RuPy(c)	4.47	1.71
ReCOPy	14.10	2.90
ReCO	0.07	0.55
TEOA	0.0021	0.0027
Py	5.19	0.87
	$(13.1^{(d)})$	$(11.0^{(d)})$

<sup>(</sup>a)  $\lambda_{\text{emission}}$ =620nm; (b)  $\lambda_{\text{emission}}$ =650nm; (c) rate constant of the  ${}^{3}$ **Ru**/ ${}^{3}$ **Ru** or  ${}^{3}$ **RuPY**/ ${}^{3}$ **RuPy** self-quenching.  $\lambda_{\text{exc}}$ =532 nm; (d) rate constant of the  ${}^{3}$ Py-moiety quenching of ( $\lambda_{\text{emission}}$ =400nm;  $\lambda_{\text{exc}}$ =355 nm). Error estimated at 5%.

#### 3.4. Molecular Insight into the preferential arrangements of the PS/Cat systems

DFT calculations were performed on the different PS/Cat systems to identify the nature and strength of interactions between their constitutive components (see computational details in ESI). The geometry of the Ru...ReCO, RuPy...ReCO, RuPy...ReCOPy...ReCOPy and RuPy...ReCOPy systems were DFT-optimized using the Becke Three-Parameter Hybrid exchange functional [46] with the Lee-Yang-Parr correlation functional [47] (B3LYP) implementing the empirical dispersion function (D3) [48] and

associated with a triple zeta basis set TZVP [49] (except for Ru and Re which were described using the LANL2DZ basis set). [50] Other computational details can be found in ESI. Their corresponding most stable conformations are reported in **Fig. S16-S22**) and their associated PS/Cat interaction energies ( $E_{int}$ ) are summarized in **Table 2**.

**Table 2.** Summary of the DFT-calculated interaction energy ( $E_{int}$ ) for the different systems (expressed in kJ mol<sup>-1</sup>). The interaction energy between the two components of each system was calculated as  $E_{int}$ =E(system)-E(complex 1)-E(complex 2) where E(system) is the total energy of the system assembling the two complexes 1 and 2 while E(complex 1) and E(complex 2) are the total energies of complex 1 and 2 taken individually, respectively. (1) and (2) correspond to two possible geometries for the ReCOPy/RuPy system, (1) being the most energetically favorable one. The determination of the interaction energy for the RuPy-RuPy system was not performed since we used a charged cation model to treat RuPy.

System	E <sub>int</sub> (kJ mol <sup>-1</sup> )
RuReCO	-156
RuPyReCO	-190
Ru…ReCOPy	-195
ReCOPyReCOPy	-261
RuPyReCOPy (1)	-295
RuPyReCOPy (2)	-227

These calculations revealed two plausible geometries for our new RuPy...ReCOPy system (labeled 1 and 2 in Table 1) as illustrated in Fig. 7a and 7b. The configuration (1) corresponds to the most energetically stable geometry resulting from the N-H/C=O hydrogen bond (d  $\approx$  1.9 Å) and the bipyridine-pyrene  $\pi$ - $\pi$ stacking (d  $\approx 2.8$  Å) between the PS and the Cat leading to the supramolecular assembly. In addition, a remarkable intramolecular alignment of pyrene functions can be clearly observed and assigned to an intramolecular  $\pi$ - $\pi$  stacking (d  $\approx$  3.4 Å). The configuration (2) is less stable and corresponds to a supramolecular assembly stabilized mainly via inter- and intra-molecular pyrene-pyrene  $\pi$ - $\pi$  stacking (d  $\approx$ 3.4 Å) as well as a N-H/C=O hydrogen bonds (d  $\approx$  1.9 Å); however, in contrast to configuration (1), the bipyridine-pyrene  $\pi$ - $\pi$  stacking interactions are absent. These configurations 1 and 2 are associated with interaction energies between RuPy and ReCOPy (-295 and/or -227 kJ.mol<sup>-1</sup> respectively) significantly higher than the value obtained between **Ru** and **ReCO** in the respective complex (-156 kJ mol<sup>-1</sup>). This result highlights that the introduction of pyrene in both Ru and ReCO leads to a substantial enhancement of the intermolecular interactions between the PS and Cat. These interactions energies are also significantly higher than the values calculated for Ru...ReCOPy (-195 kJ.mol<sup>-1</sup>) and RuPy...ReCO (-190 kJ.mol<sup>-1</sup>) (Table 2 and Fig. S17 and S18) confirming the synergistic role played by the pyrene functions present in both PS and Cat. The interaction energy for the **ReCOPy**...**ReCOPy** complex (-261 kJ.mol<sup>-1</sup>) was also found to be much higher than the value obtained for **Ru**...**ReCOPy** (-195 kJ.mol<sup>-1</sup>). This implies that the interactions between the **ReCOPy/ReCOPy** via intermolecular π-π stacking are stronger than that between the **Ru** and **ReCOPy** in the **Ru**...**ReCOPy** system (**Fig. S18**). Moreover, noting that the interaction energy for **ReCOPy**...**ReCOPy** (-261 kJ.mol<sup>-1</sup>) is lower than the value calculated for **RuPy**...**ReCOPy** (-295 kJ.mol<sup>-1</sup>) and the activity of **ReCOPy** is relatively poor without the introduction of **RuPy** (**Fig. 5b**), it comes that the 1:1 ratio maximizes the catalytic activity (**Fig. 4d**), otherwise excess of **ReCOPy** (and very likely **RuPy** as well) can group up and act similarly to the situation of **ReCOPy** only, decreasing therefore the activity. The possible electron transfer (ET) pathway for the most energetically stable **RuPy**...**ReCOPy** (1) was further explored by time-dependent DFT (TD-DFT) calculations (see ESI for computational details). These calculations did not reveal any charge transfer from the ground state of \***RuPy**. to **ReCOPy**'s (see SI: **Fig. S19** and **table S4**) as we would have expected for the associated dark reaction. Only for the second excited state, a Mulliken absolute charge difference (Δq) was obtained for the \***RuPy**. (+0.0045 |e|) and **ReCOPy** metal centers (-0.0010 |e|) with the ground state considered as a reference.

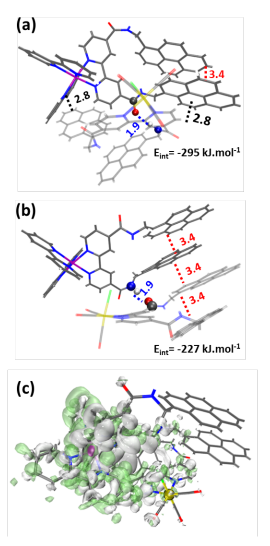


Fig. 7. DFT-calculations on the RuPy...ReCOPy system: a) the most stable configuration (1), b) plausible configuration (2) and c) charge differences for the most stable RuPy...ReCOPy (1) complex upon the excitation to the second excited state. Green and white colors represent the charge accumulation and depletion regions respectively. Color code: carbon(grey), hydrogen (white), nitrogen (blue), (green), oxygen (red), ruthenium (purple) and rhenium (yellow). The distances are reported in Å.

#### 3.5. Mechanistic investigation

After discarding the absorbance impact in the enhancement of the activity of RuPy...ReCOPy vs Ru-ReCO and based on the tests with other binary systems' combinations (Fig. 4 and S10), on the excited-state lifetimes, on the quenching rate constant and on the DFT calculations, we concluded that the higher activity is mainly attributed to a faster ET between RuPy and ReCOPy. The modelling shows two possible pathways as illustrated in Fig. 8: (i) Following a preliminary MLCT from ruthenium center to the bipyridine (bpy) and the electron then migrates from the pyrene towards the bipyridine-rhenium center to trigger the CO<sub>2</sub> photoreduction; (ii) The electron migrates from the bipyridine of RuPy towards the pyrene moieties of Re and reaches the bipyridine-rhenium center after an intermolecular transfer (Fig. 8). The interaction energy calculated by DFT between the two complexes (Table 2) allows investigating the pathway (ii) as preferential. However, pathway (i) cannot be discarded since the DFT calculations did not take into account other factors such as the solvation effect. Consequently, with the stronger interaction between RuPy and ReCOPy, both ET pathways can occur simultaneously and/or separately for ensuring a fast ET.

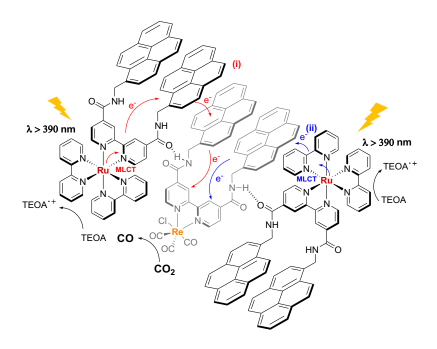


Fig. 8. The two possible electron pathways in the RuPy···ReCOPy photocatalytic systems based on the analysis of the experimental and theoretical data.

#### 4. CONCLUSION

The newly designed  $\mathbf{RuPy}$ ... $\mathbf{ReCOPy}$  supramolecular photocatalytic system was demonstrated to show an enhanced conversion efficiency of  $\mathrm{CO_2}$  to  $\mathrm{CO}$  under visible light. This supramolecular assembly is promoted by non-covalent NH/CO hydrogen bond and  $\pi$ - $\pi$  staking interactions between the catalyst and the photosensitizer. Remarkably, the calculated TOF was found three times higher than the value previously reported for the conventional  $\mathbf{Ru}$ ... $\mathbf{ReCO}$  photocatalytic system (18 h<sup>-1</sup> vs 5.3 h<sup>-1</sup>). These reaction rates and photocatalytic activity enhancement is due to an efficient electron transfer from  $\mathbf{RuPy}$  to  $\mathbf{ReCOPy}$  mediated by the pyrene moieties and the stronger interactions between  $\mathbf{RuPy}$  and  $\mathbf{ReCOPy}$  as compared to  $\mathbf{Ru}$  and  $\mathbf{ReCO}$ . The electron delocalization between the pyrene moieties and the complexes was confirmed by UV-Vis study and TR-PL showing the electron transfer pathway from ruthenium center to the pyrene moiety in  $\mathbf{RuPy}$  and from the pyrene moiety to the rhenium center in  $\mathbf{ReCOPy}$ . Additionally, the faster electron transfer from the electron donor to the  $\mathbf{RuPy}$  comparing with  $\mathbf{Ru}$  was also concluded form the quenching of the  $^3$ MLCT with TEOA. These findings shed light on the importance of the electron transfer efficiency and pathway for enhancing the performance of the photo-catalytic system for the  $\mathbf{CO_2}$  photoreduction.

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#### **Supplementary materials**

The following information are available in ESI: Ligand and complexes synthesis, NMR data, details for photocatalytic tests (in-situ reactor, calibration curve for CO quantification, FTIR spectra...), additional times resolved photoluminescence results (nanosecond emission and AT spectra, Stern-Volmert plots), energy level of HOMO and LUMO determination details and all optimized structure are presented in **Fig. S1-S22** and **Table S1-S5**.

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