

## PAPER

[View Article Online](#)  
[View Journal](#) | [View Issue](#)Cite this: *Dalton Trans.*, 2021, **50**, 14797

# DFT insights into the photocatalytic reduction of CO<sub>2</sub> to CO by Re(i) complexes: the crucial role of the triethanolamine “magic” sacrificial electron donor†

Athanasios C. Tsipis \* and Antonia A. Sarantou

The reaction mechanism for the photocatalytic reduction of CO<sub>2</sub> to CO catalyzed by the [Re(en)(CO)<sub>3</sub>Cl] complex in the presence of triethanolamine, R<sub>3</sub>N (R = CH<sub>2</sub>CH<sub>2</sub>OH) abbreviated as TEOA, in DMF solution was studied in-depth with the aid of DFT computational protocols by calculating the geometric and free energy reaction profiles for several possible reaction pathways. The reaction pathways studied start with the “real” catalytic species [Re(en)(CO)<sub>3</sub>], [Re(en)(CO)<sub>3</sub>]<sup>−</sup> and/or [Re(en)(CO)<sub>2</sub>Cl]<sup>−</sup> generated from the excited triplet **T**<sub>1</sub> state upon single and double reductive quenching by a TEOA sacrificial electron donor or photodissociation of a CO ligand. The first step in all the catalytic cycles investigated involves the capture (inserted) by the Re–OCH<sub>2</sub>CH<sub>2</sub>NR<sub>2</sub> bond forming stable intermediates. Next, successive protonations (TEOA also acts as a proton donor) lead to the release of CO either from the energy consuming 2e<sup>−</sup> reduction of [Re(en)(CO)<sub>4</sub>]<sup>+</sup> or [Re(en)(CO)<sub>2</sub>Cl]<sup>+</sup> complexes in the CO<sub>2</sub> capture pathways or from the released unstable diprotonated [R<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>OC(OH)(OH)]<sup>+</sup> species regenerating TEOA and the catalyst. The CO<sub>2</sub> insertion reaction pathway is the favorable pathway for the photocatalytic reduction of CO<sub>2</sub> → CO catalyzed by the [Re(en)(CO)<sub>3</sub>Cl] complex in the presence of TEOA manifesting its crucial role as an electron and proton donor, capturing CO<sub>2</sub> and releasing CO.

Received 1st July 2021,  
Accepted 13th September 2021

DOI: 10.1039/d1dt02188e

rsc.li/dalton

## Introduction

It is generally accepted that anthropogenic sources, such as burning of fossil fuels, are mainly responsible for the rise of atmospheric CO<sub>2</sub>, which in turn accentuates the greenhouse effect.<sup>1</sup> Many studies have been devoted to how to tackle this problem and there are continuous ongoing efforts not only to capture CO<sub>2</sub> but also to catalytically transform it into fine chemicals.<sup>2</sup> Among the various strategies to achieve this goal, the most challenging and the very attractive one is to convert CO<sub>2</sub> efficiently into useful compounds using solar light as an energy source.<sup>3</sup> Many transition metal-based complexes (Ni, Fe, Re, Cr, Ir, Mo, *etc.*) have been studied extensively for the

homogeneous electrocatalytic and photocatalytic reduction of carbon dioxide.<sup>4–16</sup>

Apaydin *et al.*<sup>7</sup> reported an excellent comprehensive overview on the homogeneous and heterogeneous CO<sub>2</sub> reduction catalyzed by organic, organometallic and bioorganic systems. Mechanistic details of the CO<sub>2</sub> reduction processes undertaken by electrochemical, bioelectrochemical and photoelectrochemical approaches are thoroughly analyzed. In parallel to the electrochemical CO<sub>2</sub> reduction, a plethora of efforts were focused on the photocatalytic CO<sub>2</sub> reduction, which relied on systems comprising a light harvesting unit consisting of a photosensitizer (PS) compound and two catalytic sites.<sup>8–14</sup> In the oxidation site, a donor provides an electron e<sup>−</sup> to the PS after its excitation to the triplet excited state (<sup>3</sup>MLCT) which is subsequently reductively quenched by the reduction site and finally, the e<sup>−</sup> is transferred to CO<sub>2</sub>. However, in many cases, the PS acts not only as a photosensitizer but also as a reduction site as well.

Among the mononuclear transition metal complexes developed so far for electro- and photocatalytic CO<sub>2</sub> reduction, polypyridyl transition metal complexes constitute the class of molecular catalysts employed in the reduction of CO<sub>2</sub> to CO. An excellent overview of the CO<sub>2</sub> reduction catalyzed by poly-

Department of Chemistry, University of Ioannina, Ioannina 45110, Greece.

E-mail: attsipis@uoi.gr

† Electronic supplementary information (ESI) available: Reaction steps for the photocatalytic reduction of CO<sub>2</sub> to CO catalyzed by the [(en)(CO)<sub>3</sub>ReCl] catalyst starting with the [(en)(CO)<sub>3</sub>Re] intermediate resulted upon one electron reduction of the **T**<sub>1</sub> state by TEOA (Fig. S1). Cartesian coordinates of the reactants, intermediates and products involved in the catalytic cycles (Table S1). See DOI: 10.1039/d1dt02188e

pyridyl transition metal complexes has recently been published by Fontecave's group,<sup>8</sup> presenting and thoroughly analyzing the proposed catalytic cycles for the CO<sub>2</sub> reduction by Re(bpy)CO<sub>3</sub>(X) investigated by both experimental and computational approaches. Generally, mechanistic studies indicated that, initially a two electron reduction of the Re(bpy)CO<sub>3</sub>(X) catalyst affords the anionic five-coordinated 17e<sup>-</sup> [Re(bpy)(CO)<sub>3</sub>]<sup>-</sup> complex which captures CO<sub>2</sub> forming the [Re(bpy)(CO)<sub>3</sub>(CO<sub>2</sub>)]<sup>-</sup> intermediate, which upon protonation yields the Re(bpy)(CO)<sub>3</sub>(CO<sub>2</sub>H) intermediate which upon second protonation and additional electron reduction is converted into [Re(bpy)(CO)<sub>4</sub>]<sup>-</sup>. However, a question still remains concerning the release of the CO product from the [Re(bpy)(CO)<sub>4</sub>]<sup>-</sup> complex. More recently, Ishitani and co-workers<sup>11</sup> presented an excellent discussion of all the reaction mechanisms proposed for the photochemical CO<sub>2</sub> reduction catalyzed by Re(I) and Ru(II) complexes. The authors stated that no one from the numerous mechanisms proposed is a universal mechanism. This review article is recommended to the readers and researchers in the field. On the other hand, Cramer's group reported mechanistic details on the proton-dependent electrocatalytic reduction of CO<sub>2</sub> to CO by *fac*-Re(bpy)(CO)<sub>3</sub>Cl using first principles quantum chemistry.<sup>13</sup> The Cramer's group also compared the complete electrocatalytic cycles of CO<sub>2</sub> to CO reduction catalyzed by *fac*-Re(bpy)(CO)<sub>3</sub>Cl and *fac*-Mn(bpy)(CO)<sub>3</sub>Cl catalysts.<sup>14</sup>

The photo-induced reduction of CO<sub>2</sub> to CO in the acetonitrile/water/triethylamine solution in the presence of a [Ru(2,2'-bipyridine)<sub>3</sub>]<sup>2+</sup>/CO<sub>2</sub><sup>+</sup> system was first reported by Lehn and Ziessel.<sup>17</sup> In subsequent publications, Lehn's group<sup>18,19</sup> showed that the most selective reduction catalysts to produce CO from CO<sub>2</sub> are Re(I) octahedral complexes with the general formula [Re(bpy)(CO)<sub>3</sub>X] (X = Cl, Br). The proposed catalytic cycle for the photoreduction of CO<sub>2</sub>, catalyzed by the (bpy)Re(CO)<sub>3</sub>X complexes, comprises excitation to the <sup>3</sup>MLCT state and reduction of the Re(I) catalyst using triethanolamine (TEOA) as an electron donor yielding the one electron reduced Re(I) complex (OER-species) that loses the X<sup>-</sup> ligand forming an unstable 17e<sup>-</sup> species. Next, the 17e<sup>-</sup> Re(I) complex could capture CO<sub>2</sub> *via* coordination to the rhenium metal center.<sup>17-19</sup> However, the mechanism of conversion of CO<sub>2</sub> to CO has not yet been fully understood and there are some points that are still under debate and need to be clarified.

The mechanism of the photoinduced reduction of CO<sub>2</sub> to CO in a TEOA/DMF/[ReBr(CO)<sub>3</sub>(bpy)] system has been investigated by Kutal *et al.*<sup>20,21</sup> Reductive quenching of the photoexcited [ReBr(CO)<sub>3</sub>(bpy)] complex by TEOA affords the reduced [Re'Br(CO)<sub>3</sub>(bpy<sup>-</sup>)] species which can be viewed as a Re' center bound to a 2,2'-bipyridine radical anion. The TEOA<sup>+</sup> generated can rapidly abstract a hydrogen atom from another TEOA molecule to produce the strong reducing radical, TEOA<sup>•</sup>. Next, the 19e<sup>-</sup> Re species activates CO<sub>2</sub> for reduction to CO, but the composition, structure, or subsequent reactivity of any intermediates formed is unknown.

Although several studies<sup>22-29</sup> point towards the existence of the 17e<sup>-</sup> reactive species, the next step involving capturing CO<sub>2</sub> by the catalytic system is unclear. Kubiak and co-workers<sup>22-25</sup>

investigated the catalytic activity of the Re(bpy)(CO)<sub>3</sub>X complexes (bpy = 4,4'-dicarboxyl-2,2'-bipyridine, 2,2'-bipyridine, 4,4'-dimethyl-2,2'-bipyridine, 4,4'-di-*tert*-butyl-2,2'-bipyridine, and 4,4'-dimethoxy-2,2'-bipyridine) and found that Re(bpy-*t*Bu)(CO)<sub>3</sub>Cl has the most significant catalytic activity for the reduction of CO<sub>2</sub> to CO. DFT calculations showed that the geometry of the [Re(bpy-*t*Bu)(CO)<sub>3</sub>(CO<sub>2</sub>)]<sup>-</sup> doubly reduced species converges only upon inclusion of a cation (H, Li, Na, or K). Based on these studies, the authors proposed a catalytic cycle that involves a two-electron reduction of the [Re(bpy-R)(CO)<sub>3</sub>X] (R = H, Me, *t*Bu, X = halogen, OTf) complex which upon dissociation of the X ligand yields the anionic [Re(bpy-R)(CO)<sub>3</sub>]<sup>-</sup> species. Several [Re(bpy-R)(CO)<sub>3</sub>]<sup>-</sup> anions were isolated and fully characterized.<sup>25</sup> Next, the CO<sub>2</sub> molecule is coordinated to the Re metal center yielding a carboxylate [Re(bpy-R)(CO)<sub>3</sub>(CO<sub>2</sub>)]<sup>-</sup> intermediate, which upon protonation forms the carboxylato [Re(bpy-R)(CO)<sub>3</sub>(CO<sub>2</sub>H)] complex. Further protonation of the [Re(bpy-R)(CO)<sub>3</sub>(CO<sub>2</sub>H)] complex releases H<sub>2</sub>O and produces the cationic tetracarbonyl species. The release of CO from the [Re(bpy-R)(CO)<sub>4</sub>]<sup>+</sup> complex is catalysed by electron transfer.

On the other hand, Muckerman *et al.*<sup>26</sup> in a prominent theoretical study precluded direct coordination of the Re metal center and instead proposed that the catalytic cycle proceeds *via* the formation of a Re dimeric intermediate species. In the latter, CO<sub>2</sub> acts as a bridging ligand between the two Re metal centers, though the formation of the carboxylate dimeric species is entropically unfavorable. Ishitani *et al.*<sup>27</sup> proposed an alternative route by revealing the crucial role of TEOA in the CO<sub>2</sub> capture by the Re(I) catalyst. The authors proposed a catalytic cycle for the photocatalytic CO<sub>2</sub> reduction catalyzed by the [Re(bpy)(CO)<sub>3</sub>(NCS)] complex that involves excitation of the S<sub>0</sub> ground state of the [Re(bpy)(CO)<sub>3</sub>(NCS)] complex to the triplet <sup>3</sup>MLCT state, which subsequently is reductively quenched by TEOA, giving the one electron reduced (OER) [Re(bpy)(CO)<sub>3</sub>(NCS)]<sup>-</sup> species. The dissociation of the SCN<sup>-</sup> ligand from the OER yields a 17e<sup>-</sup> five-coordinated intermediate which reacts with CO<sub>2</sub> to give the CO<sub>2</sub> adduct(s). Further electron donation to the OER results in the release of CO and the [Re(bpy)(CO)<sub>3</sub>]<sup>+</sup> complex and regenerates the catalyst. Later Ishitani's group<sup>28</sup> performing spectrochemical and cyclic voltammetry measurements on adding TEOA to a solution of *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(CH<sub>3</sub>CN)]PF<sub>6</sub> dissolved in pure DMF provided evidence for the existence of the *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(OCH<sub>2</sub>CH<sub>2</sub>NR<sub>2</sub>)] (R = CH<sub>2</sub>CH<sub>2</sub>OH) intermediate in solution. A Re(I) intermediate with the CO<sub>2</sub> inserted between the Re(I) metal center and the TEOA anionic ligand was isolated and fully characterized.

The photocatalytic activity of transition metal-based catalysts towards the reduction of CO<sub>2</sub> to CO depends upon the structure of the complexes. Attempts to address this issue have already been made. For example, Kurz *et al.*<sup>29</sup> performed an experimental structure-activity study of [(diimine)(CO)<sub>3</sub>ReX] complexes with respect to their ability to act as photocatalysts for the reduction of CO<sub>2</sub> to CO. Also, Ishida *et al.*<sup>30</sup> studied the effect of ligand L on the photocatalytic activity for CO<sub>2</sub> conver-



sion in a series of *trans*-(Cl)-[Ru(L)(CO)<sub>2</sub>Cl<sub>2</sub>] complexes, revealing the significance of their reduction potential,  $E_p$ . The latter is modulated by the Lowest Unoccupied Molecular Orbital (LUMO) eigenvalue which in turn depends on the nature of L.

In this regard, the main goal that we pursued in this work was to scrutinize the mechanistic details of the photocatalytic conversion of CO<sub>2</sub> to CO by the octahedral [(en)(CO)<sub>3</sub>ReCl] (en = ethylenediamine) complex selected as a model complex, employing DFT methods. Our efforts were directed towards the elucidation of the mechanism of the reaction step related to CO<sub>2</sub> capture and CO release in the reduction of CO<sub>2</sub> to CO catalyzed by Re(I) complexes in DMF/TEOA solutions.

## Computational methods

All stationary points (reactants, transition states, and products) located on the potential energy surfaces (PES) were fully optimized at the wB97XD/Def2-TZVP level of theory as implemented in the Gaussian16 suite of programs.<sup>31</sup> The wB97XD functional<sup>32–35</sup> containing empirical dispersion terms and long-range corrections provides good descriptions of the reaction profiles, including geometries, heats of reaction, and barrier heights.<sup>36</sup> Analytical frequencies were calculated at the same level of theory, and the nature of the stationary points was determined in each case according to the number of the negative eigenvalues of the Hessian matrix. Gibbs free energies were calculated at 298.150 Kelvin and 1 Atm. Solvent effects were accounted for by means of the Polarizable Continuum Model (PCM) with the integral equation formalism variant (IEF-PCM) being the default self-consistent reaction field (SCRF) method.<sup>37</sup> DMF was used as the solvent. Natural Bond Orbital (NBO) population analysis was performed using Weinhold's methodology.<sup>38,39</sup> Time-dependent density functional theory (TD-DFT)<sup>40–42</sup> calculations were performed on the equilibrium ground state geometries in DMF solution employing the wB97XD/Def2-TZVP/PCM computational protocol.

## Results and discussion

### Computation-based catalytic cycles

The photocatalytic CO<sub>2</sub> → CO transformation catalyzed by the selected [(en)(CO)<sub>3</sub>ReCl] model catalyst proceeds through three main reaction steps: (i) step A1 corresponds to the photo-excitation of the [(en)(CO)<sub>3</sub>ReCl] catalyst accompanied by one electron reduction of the triplet excited state **T**<sub>1</sub> by TEOA, the so-called “magic” sacrificial electron donor,<sup>28</sup> yielding very reactive coordinatively unsaturated, 17e<sup>−</sup> intermediates that are the catalytically active species initiating step B1. (ii) Step B1 involves the interaction of the oxidized TEOA<sup>+</sup> species formulated as [R<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>OH]<sup>+</sup> (R = CH<sub>2</sub>CH<sub>2</sub>OH) with the Re metal center yielding [(en)(CO)<sub>3</sub>Re(OCH<sub>2</sub>CH<sub>2</sub>NR<sub>2</sub>)] intermediates capable of capturing CO<sub>2</sub> by insertion into the Re–O bond. Step B1 has thoroughly been investigated both experi-

mentally and theoretically by Ishitani's group<sup>28</sup> employing the *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(CH<sub>3</sub>CN)]PF<sub>6</sub> catalyst. (iii) Step C starts with the [R<sub>2</sub>NCH<sub>2</sub>CH<sub>2</sub>C(O)OH]<sup>+</sup> species released upon protonation of the [(en)(CO)<sub>3</sub>ReO(O)(COCH<sub>2</sub>CH<sub>2</sub>NR<sub>2</sub>)] intermediate and its conversion upon further protonation to the desired CO molecule, H<sub>2</sub>O and regenerating TEOA. Step C manifests the crucial role of TEOA in the photocatalytic reduction of CO<sub>2</sub> → CO, thus clarifying, the “magic” sacrificial electron donor features of TEOA.<sup>28</sup>

The reaction steps for the photocatalytic reduction of CO<sub>2</sub> to CO catalyzed by the [(en)(CO)<sub>3</sub>ReCl] (en = ethylenediamine) catalyst, calculated at the wB97XD/def2-TZVP level, in a DMF solution are depicted schematically in the ESI (Fig. S1†). The alternative reaction steps B1' and B1'' starting with the coordination of CO<sub>2</sub> to the Re metal center of **ImA1** and/or **ImA1'** (one electron reduced **ImA1** species) intermediates yielding the [Re(en)(CO)<sub>3</sub>(CO<sub>2</sub>)] and/or [Re(en)(CO)<sub>3</sub>(CO<sub>2</sub>)]<sup>−</sup> intermediates, which upon successive protonations afford the tetracarbonyl [Re(en)(CO)<sub>4</sub>]<sup>2+</sup> and [Re(en)(CO)<sub>4</sub>]<sup>+</sup> complexes respectively will be discussed separately and compared with step B1. We also investigated alternative reaction pathways starting with a five-coordinated [(en)(CO)<sub>2</sub>ReCl]<sup>−</sup> intermediate that resulted from the dissociation of one of the equatorial CO ligands of the excited **T**<sub>1</sub> triplet state (step A2) which will be discussed later on.

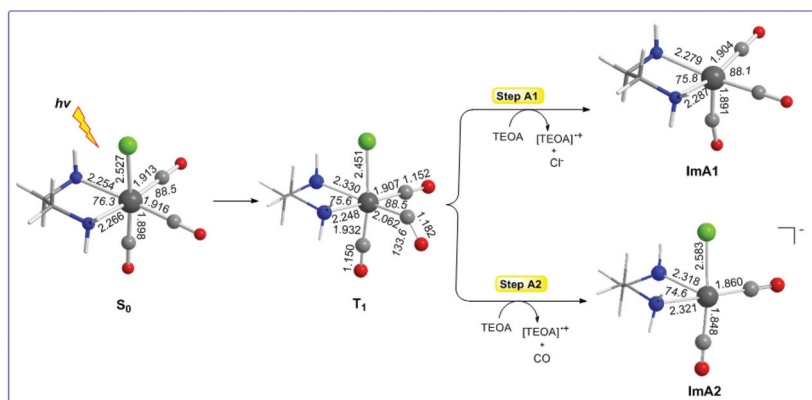
### Reaction steps A1 and A2: photophysics of the [(en)(CO)<sub>3</sub>ReCl] catalyst

Let us examine first steps A1 and A2 concerning the photo-excitation of the [(en)(CO)<sub>3</sub>ReCl] catalyst. Since excitation of the [(en)(CO)<sub>3</sub>ReCl] complex is a prerequisite in order to initiate the photocatalytic reduction of CO<sub>2</sub> to CO, we set out to study the photophysical properties by means of Time Dependent DFT (TDDFT) electronic structure calculations. The simulated absorption spectrum of the [(en)(CO)<sub>3</sub>ReCl] complex calculated by the TDDFT/wB97XD/Def2-TZVP/PCM computational protocol shows absorption bands spanning between 200 and 300 nm with the most intense among them peaking at 211, 231 and 273 nm. The peak at 211 nm exhibits a complex nature and could be assigned as Metal to Ligand/Ligand to Ligand Charge Transfer (MLCT/LL'CT). The same assignment holds also true for the electronic transitions absorbing at 231 and 273 nm as well.

The equilibrium geometries of the [(en)(CO)<sub>3</sub>ReCl] complex in the ground **S**<sub>0</sub> and excited triplet **T**<sub>1</sub> states along with the catalytically active five-coordinated [(en)(CO)<sub>3</sub>Re] and [(en)(CO)<sub>2</sub>ReCl] species formed in photo-physical steps A1 and A2 respectively are shown in Fig. 1. Calculations predicted that the *fac*-[(en)(CO)<sub>2</sub>ReCl] isomer is more stable than the *mer*-[(en)(CO)<sub>2</sub>ReCl] isomer in both the **S**<sub>0</sub> and **T**<sub>1</sub> states by 24.1 and 6.8 kcal mol<sup>−1</sup> respectively.

It can be seen that one electron transfer from TEOA to the **T**<sub>1</sub> state of [(en)(CO)<sub>3</sub>ReCl] promotes the cleavage of either the Re–Cl or the Re–CO bond in the reduced 19e<sup>−</sup> transient **OER** species formed, yielding the very reactive 17e<sup>−</sup> [(en)(CO)<sub>3</sub>Re], **ImA1** and [(en)(CO)<sub>2</sub>ReCl]<sup>−</sup>, **ImA2** intermediates respectively.





**Fig. 1** Equilibrium geometries of the  $[(en)(CO)_3ReCl]$  complex in the ground  $S_0$  and excited triplet  $T_1$  states along with the catalytically active five-coordinated  $[(en)(CO)_3Re]$ , **ImA1** and  $[(en)(CO)_2ReCl]$ , **ImA2** species formed in photo-physical steps A1 and A2 respectively calculated at the wB97XD/Def2-TZVP level of theory in a DMF solution.

Noteworthy in  $T_1$  one of the equatorial CO ligands forms a weaker Re–CO bond (Re–CO bond length of 2.062 Å) compared to the Re–CO bonds of the other two CO ligands in the equatorial and axial coordination sites with bond lengths of 1.907 and 1.932 Å respectively. Interestingly the Re–C–O bond angle of the weakly bonded CO ligand is 133.6°. Scheme 1 shows the 3D plots of the frontier molecular orbitals (FMOs) and the spin density distribution of  $T_1$ , along with the estimated Wiberg Bond Indices (WBOs) for the Re–Cl and Re–CO bonds in  $T_1$  (the entry in the right side of Scheme 1).

It can be seen that the Single Occupied Molecular Orbital (SOMO) in  $T_1$  is mainly localized on the weakly bonded CO ligand. The SOMO is composed of 19.1% Re spd, 45.2% C sp hybrid orbitals and 18.9% O p orbitals. The Lowest Unoccupied Molecular Orbital (LUMO) is composed of 25.8% Re spd and 24.0% C sp hybrid orbitals. The spin density is distributed on the Re–CO bond region, 1.078|e| on Re and 0.716 on the CO ligand. The estimated WBI(Re–CO) shows also their bond strengths, being 1.187 for the weaker bonded CO ligand and 1.720 and 1.576 for the CO ligands coordinated at the equatorial and axial coordination sites respectively.

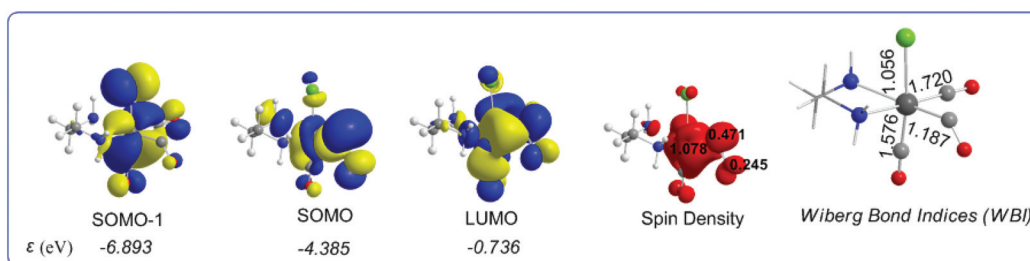
### Reaction step B1: CO<sub>2</sub> insertion

The geometric and free energy reaction profiles calculated for step B1 are shown in Fig. 2.

Initially, a DMF molecule interacts with the very reactive  $17e^-$  five coordinate intermediate **ImA1**, with the estimated interaction energy (IE) being  $-7.4 \text{ kcal mol}^{-1}$  forming a loose association through an endergonic process ( $\Delta G = 4.5 \text{ kcal mol}^{-1}$ ). The **ImA1**...DMF interactions correspond to non-covalent interactions clearly shown in Scheme 2.

Next, DMF is substituted by an  $OCH_2CH_2NR_2$  ligand yielding an **ImB1'** intermediate, in line with the experimental observations by Ishitani's group<sup>28</sup> that upon adding TEOA to a solution of the  $fac-[Re^I(bpy)(CO)_3(DMF)]^+$  complex (**Im\_DMF**), a 1 : 2 thermodynamically equilibrated mixture of **Im\_DMF** with the  $fac-[Re^I(bpy)(CO)_3(OCH_2CH_2NR_2)]$  (**ImB**) intermediate is obtained. The equilibrium between **Im\_DMF** and **ImB** intermediates was further verified by IR spectroscopy.<sup>11</sup> The substitution reaction is exergonic ( $\Delta G = -68.7 \text{ kcal mol}^{-1}$ ).

Natural Bond Orbital (NBO) population analysis revealed that the  $OCH_2CH_2NR_2$  ligand is coordinated to the Re metal centre of the  $[(en)(CO)_3Re]$  catalytic species, **ImA1**, forming a relatively weak Re–O bond with an estimated WBI(Re–O) of 0.550. Note that both the Re metal centre and the O donor atom of the coordinated  $OCH_2CH_2NR_2$  ligand acquire negative natural atomic charges of  $-0.592$  and  $-0.779|e|$  respectively. The bonding  $\sigma(Re-O)$  NBO is constructed from the interaction of  $sp^{2.69}d^{2.88}$  hybrid orbitals (40.8% p and 43.7% d character)



**Scheme 1** 3D plots of the frontier molecular orbitals (FMOs), spin density distribution and Wiberg Bond Indices (WBI) of  $T_1$ , calculated at the wB97XD/Def2-TZVP level of theory.





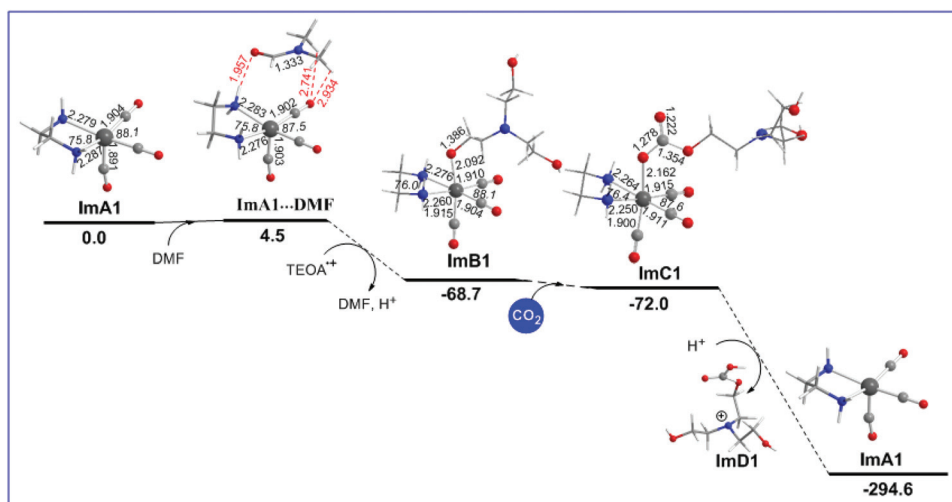
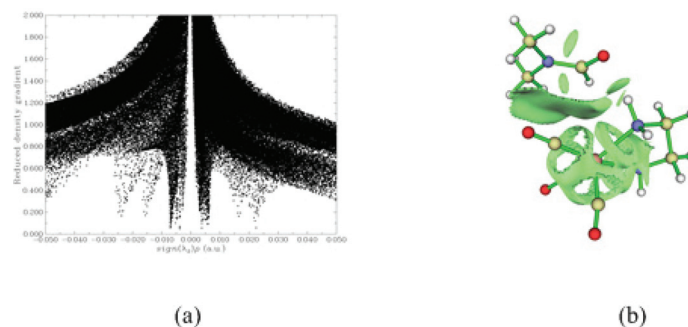


Fig. 2 Geometric and free energy,  $\Delta G$  (in  $\text{kcal mol}^{-1}$ ), reaction profiles for step B of the reduction of  $\text{CO}_2$  to CO catalyzed by the  $[(\text{en})(\text{CO})_3\text{ReCl}]$  catalyst calculated at the wB97XD/Def2-TZVP level in the DMF solution. The sum of  $G_{\text{ImA1}} + G_{\text{DMF}}$  at an infinite distance was considered to be 0.00.



Scheme 2 Scatter graphs of the RDG vs.  $\text{sign}(\lambda_2)\rho(r)$  correlations (a) along with the 3D RDG isosurfaces (isosurface = 0.70 au) for the **ImA1...DMF** intermediate calculated at the wB97XD/Def2-TZVP level of theory.

of Re with an  $\text{sp}^{1.82}$  hybrid (64.5% p-character) on the oxygen donor atom and is described as  $\sigma(\text{Re}-\text{O}) = 0.395h_{\text{Re}} + 0.919h_{\text{O}}$ . The occupancy of the  $\sigma(\text{Re}-\text{O})$  NBO is  $1.939|e|$ .

Next, a  $\text{CO}_2$  molecule is inserted into the  $\text{Re}-\text{OCH}_2\text{CH}_2\text{NR}_2$  bond transforming the  $\text{OCH}_2\text{CH}_2\text{NR}_2$  to the  $\text{OC}(\text{O})\text{OCH}_2\text{CH}_2\text{NR}_2$  ligand yielding intermediate **ImC1**. The insertion process is slightly exergonic by only  $-3.3 \text{ kcal mol}^{-1}$  (Fig. 2). **ImC1** could be subjected to protonations at the O atoms of the coordinated  $\text{OC}(\text{O})\text{OCH}_2\text{CH}_2\text{NR}_2$  ligand marked as  $\text{O}_a$  and  $\text{O}_b$  protonation sites (Fig. S1†). The natural atomic charges at the  $\text{O}_a$  and  $\text{O}_b$  atoms are  $-0.759$  and  $-0.708|e|$  respectively. The protonation either at  $\text{O}_a$  or  $\text{O}_b$  of the coordinated  $[\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}-\text{C}(\text{O})\text{O}]$  ligand forces the rupture of the  $\text{Re}-\text{O}$  bond which yields **ImD1** formulated as  $[\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}-\text{C}(\text{O})\text{OH}]^+$  and regenerates the five coordinated  $17e^-$  **ImA1** catalytic species, thus closing the catalytic cycle. The protonation accompanied by ligand dissociation corresponds to a strongly exergonic process ( $\Delta G = -222.6 \text{ kcal mol}^{-1}$ ).

### Reaction step C

**ImD** is the species initiating reaction step C that delivers the desired CO product. The geometric and free energy reaction profiles calculated for step C are shown in Fig. 3. As in the case of **ImC1** (*vide supra*), there are two possible protonation sites in **ImD** (marked as a and b in Fig. S1†). The natural atomic charges on the  $\text{O}_a$  and  $\text{O}_b$  atoms of **ImD** are  $-0.668$  and  $-0.659|e|$  respectively. Thus, the protonation at the  $\text{O}_a$  site is the most favoured one yielding **ImF** which upon further protonation and one electron reduction delivers the CO product and regenerates TEOA. The **ImD**  $\rightarrow$  **ImF** transformation is strongly exergonic ( $\Delta G = -107.5 \text{ kcal mol}^{-1}$ ). **ImF** can also be obtained upon protonation at the  $\text{O}_b$  site in **ImD** forming a transition state **TS** ( $\nu_i = -1888 \text{ cm}^{-1}$ ) surmounting an activation barrier of  $41.1 \text{ kcal mol}^{-1}$  (Fig. 1). The normal coordinate vectors (arrows) of the vibrational modes corresponding to the imaginary frequency of **TS** show that the dominant motions reflect the structural changes



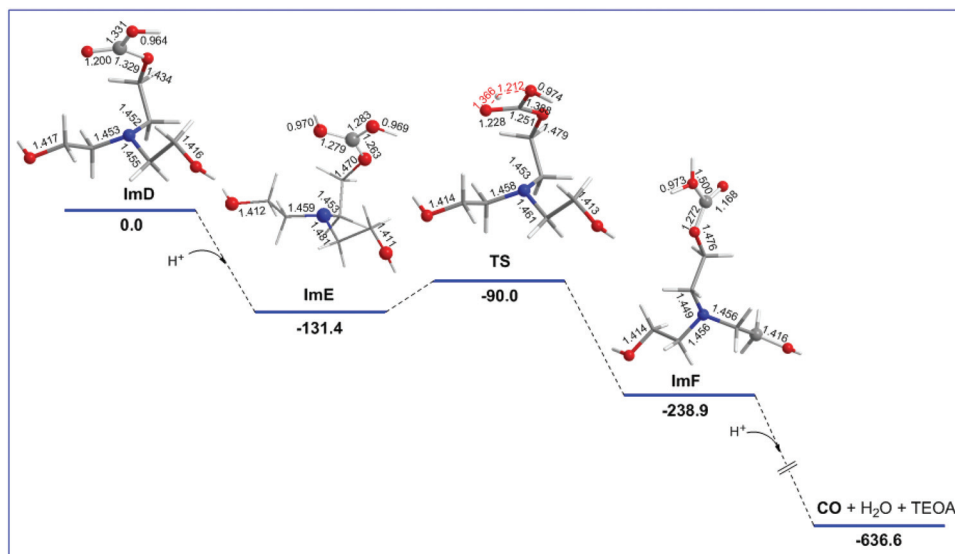


Fig. 3 Geometric and free energy,  $\Delta G$  (in  $\text{kcal mol}^{-1}$ ), reaction profiles for step C of the reduction of  $\text{CO}_2$  to CO catalyzed by the  $[(\text{en})(\text{CO})_3\text{ReCl}]$  catalyst calculated at the wB97XD/Def2-TZVP level in the DMF solution.

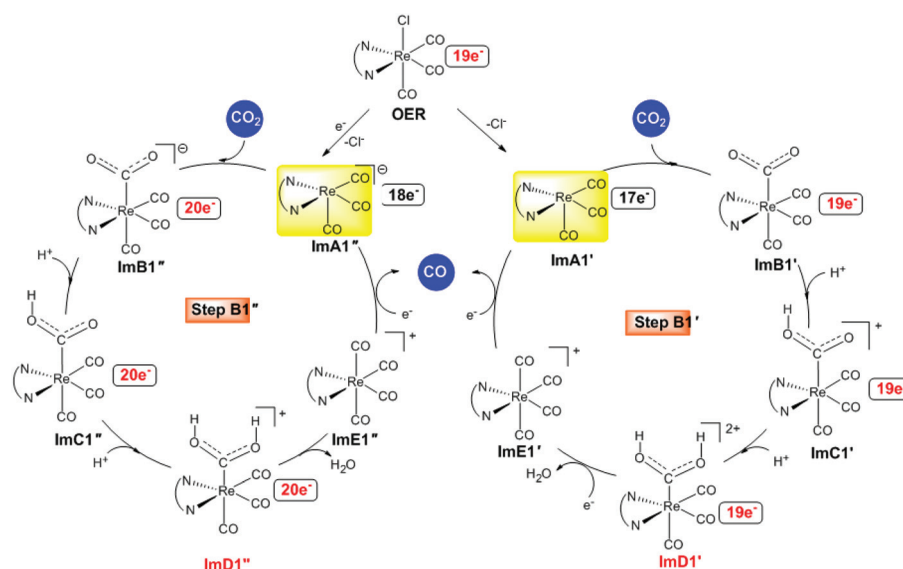


Fig. 4 Proposed catalytic cycles for steps B1' and B1'' calculated at the wB97XD/Def2-TZVP level in the DMF solution.

leading to the formation of **ImF**. The relatively high activation barrier indicates that the **ImD**  $\rightarrow$  **TS**  $\rightarrow$  **ImF** is not a favoured pathway. Noteworthy reaction step C is a key step that manifests the crucial role of the TEOA sacrificial electron donor in the photocatalytic reduction of  $\text{CO}_2$  to CO catalyzed by  $\text{Re}(\text{I})$  complexes.

#### Reaction steps B1' and B1'': $\text{CO}_2$ capture

Alternative reaction pathways (steps B1' and B1'') involving direct capture of  $\text{CO}_2$  by the active **ImA1'** catalytic species (step B1') or its one electron reduced **ImA1''** species (step B1'') yield-

ing the  $[\text{Re}(\text{en})(\text{CO})_3(\text{CO}_2)]$  and  $[\text{Re}(\text{en})(\text{CO})_3(\text{CO}_2)]^-$  intermediates, respectively have also been explored by DFT methods. The proposed catalytic cycles for steps B1' and B1'' are shown in Fig. 4. The geometric and free energy reaction profiles calculated for steps B1' and B1'' are given in Fig. 5 and 6 respectively.

After excitation of the  $S_0$  ground state of the  $[\text{Re}(\text{en})(\text{CO})_3\text{Cl}]$  complex to the triplet  $T_1$  state, the latter could undergo one and two electron reductions by TEOA accompanied by  $\text{Cl}^-$  dissociation affording the catalytically active  $17e^-$  **ImA1'** and the anionic  $18e^-$  **ImA1''** intermediates which initiate reaction steps B1' and B1'' respectively (Fig. 4).



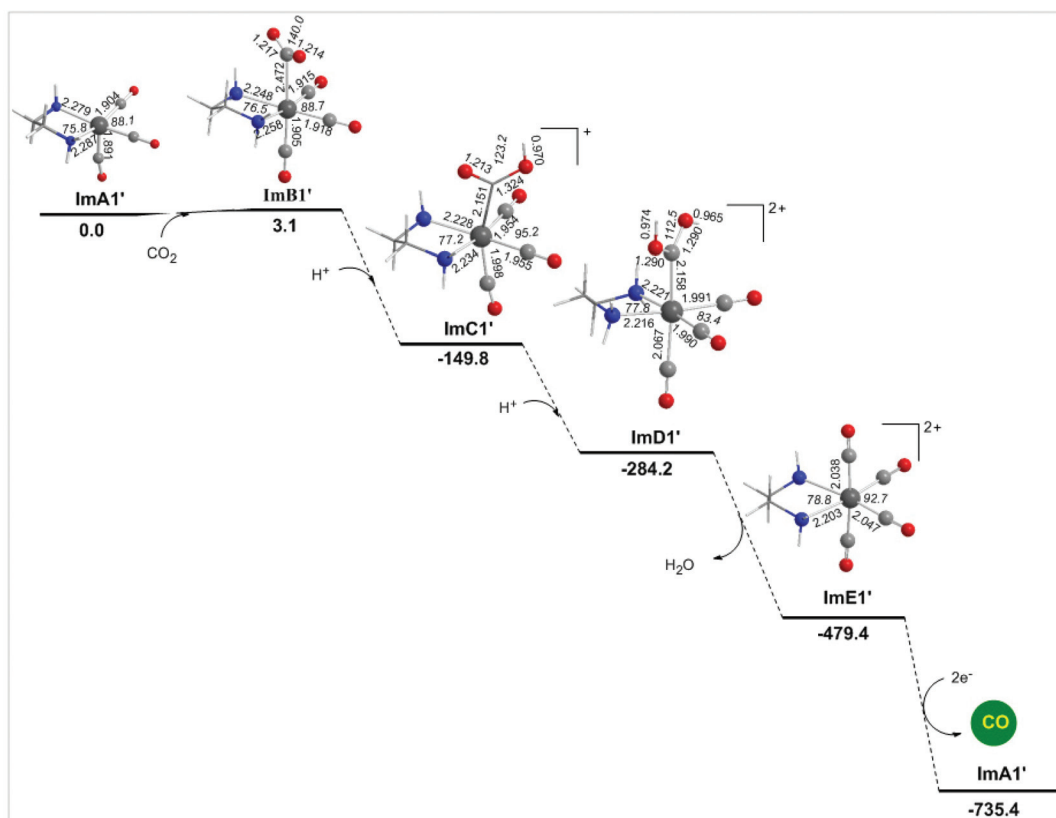


Fig. 5 Geometric and free energy,  $\Delta G$  (in kcal mol<sup>-1</sup>), reaction profiles for step B1' of the reduction of CO<sub>2</sub> to CO catalyzed by the [(en)(CO)<sub>3</sub>ReCl] catalyst calculated at the wB97XD/Def2-TZVP level in the DMF solution.

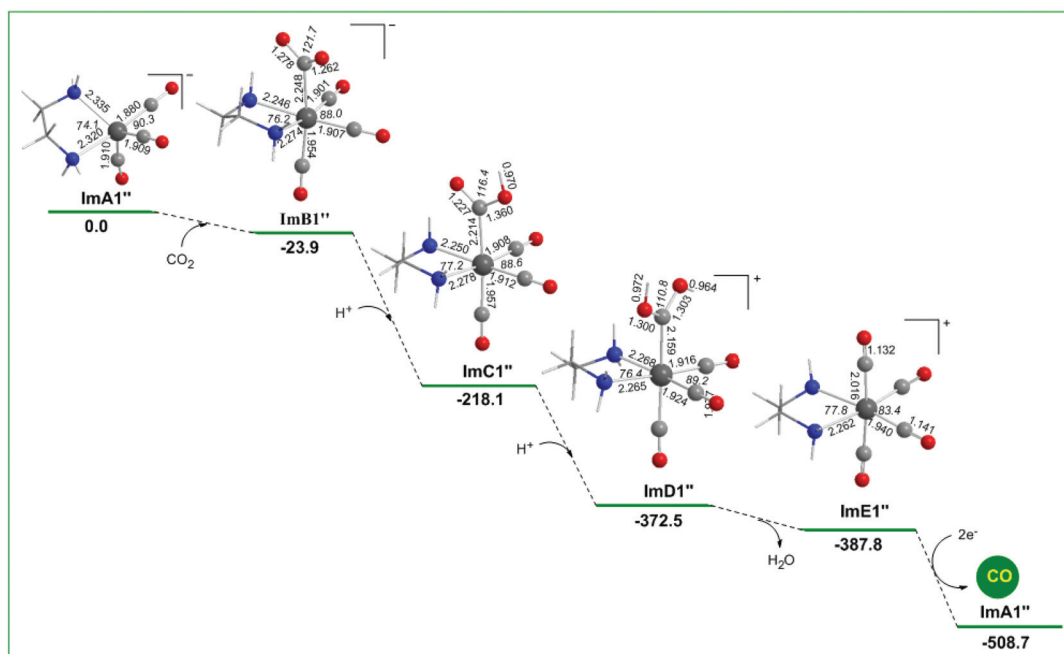


Fig. 6 Geometric and free energy,  $\Delta G$  (in kcal mol<sup>-1</sup>), reaction profiles for step B1'' of the reduction of CO<sub>2</sub> to CO catalyzed by the [(en)(CO)<sub>3</sub>ReCl] catalyst calculated at the wB97XD/Def2-TZVP level in the DMF solution.



According to NBO population analysis, the Re metal centers in **ImA1'** and **ImA1''** intermediates acquire negative natural atomic charges of  $-0.752$  and  $-1.174|e|$  respectively. Noteworthy the Re metal center in **ImA1'** and **ImA1''** is a nucleophilic center and therefore could not be attacked by a nucleophile. In this regard, direct coordination of CO<sub>2</sub> (CO<sub>2</sub> capture) to the Re metal center of **ImA1'** and **ImA1''** intermediates seems to be highly unlikely in line with the experimental observations reported by Ishitani's group<sup>28</sup> that no reaction occurs when CO<sub>2</sub> is added to a solution of the *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(DMF)]<sup>+</sup> complex, while the capture of CO<sub>2</sub> was observed when bubbling CO<sub>2</sub> into an equilibrium mixture of *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(DMF)]<sup>+</sup>/*fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>(OCH<sub>2</sub>CH<sub>2</sub>NR<sub>2</sub>)] (bpy = 2,2'-bipyridine) complexes in solution by <sup>1</sup>H and <sup>13</sup>C NMR spectroscopy yielding the *fac*-[Re<sup>I</sup>(bpy)(CO)<sub>3</sub>{R<sub>2</sub>N-CH<sub>2</sub>CH<sub>2</sub>O-COO}] complex which was isolated and fully characterized.

Scheme 3 shows the 3D plots of the frontier molecular orbitals (FMOs) and spin density distributions of **ImA1'** and **ImA1''** catalytic species.

The capture of CO<sub>2</sub> by **ImA1'** in the reaction step B1' yielding **ImB1'** (Fig. 6) is slightly endergonic ( $3.1 \text{ kcal mol}^{-1}$ ), while the estimated interaction energy is  $7.0 \text{ kcal mol}^{-1}$ . According to the NBO population analysis, the CO<sub>2</sub> molecule is loosely associated with the Re metal centre of the **ImA1'** catalytic species, with the Re–CO<sub>2</sub> bond having a WBI(Re–CO<sub>2</sub>) of only 0.348. Note that the Re metal centre acquires a negative natural atomic charge of  $-0.808|e|$  and the C atom of the coordinated CO<sub>2</sub> a positive natural atomic charge of  $0.755|e|$ . The side-on orientation of the CO<sub>2</sub> molecule in the **ImB1'** intermediate indicates that non-covalent (dipole–dipole and electrostatic) interactions predominate in the coordination of CO<sub>2</sub> to the **ImA1'** catalytic species. The weak covalent interactions are mirrored on the bonding  $\sigma(\text{Re}–\text{CO}_2)$  NBO which is constructed from the interaction of the  $\text{sp}^{3.52}\text{d}^{2.74}$  hybrid orbitals (48.4% p and 37.7% d character) of Re with an  $\text{sp}^{2.30}$  hybrid (69.6% p-character) on the carbon donor atom of the CO<sub>2</sub> ligand and described as  $\sigma(\text{Re}–\text{CO}_2) = 0.605h_{\text{Re}} + 0.797h_{\text{C}}$ . The occupancy or the  $\sigma(\text{Re}–\text{CO}_2)$  NBO is  $0.947|e|$ .

Next the successive protonations of the coordinated CO<sub>2</sub> yielding **ImC1'** and **ImD1'** intermediates are exergonic processes, and the estimated  $\Delta G$  values are  $-152.9$  and  $-134.6 \text{ kcal mol}^{-1}$  respectively. **ImD1'** releasing H<sub>2</sub>O affords the dicationic [Re(en)(CO)<sub>4</sub>]<sup>2+</sup> complex **ImE1'** which upon 2e<sup>−</sup> reduction releases the desired CO product and regenerates the catalytically active **ImA1'** species.

In reaction step B1'' (Fig. 6), the capture of CO<sub>2</sub> by **ImA1''** yielding **ImB1''** is exergonic ( $-23.9 \text{ kcal mol}^{-1}$ ), indicative of the stronger non-covalent and covalent interactions between **ImA1''** and CO<sub>2</sub> (with an interaction energy of  $23.9 \text{ kcal mol}^{-1}$ ). The NBO population analysis revealed that the CO<sub>2</sub> molecule is coordinated to the Re metal centre of the **ImA1''** catalytic species, forming a stronger nearly single Re–CO<sub>2</sub> bond with a WBI(Re–CO<sub>2</sub>) of 0.755. The Re metal centre acquires a negative natural atomic charge of  $-1.016|e|$  and the C atom of the coordinated CO<sub>2</sub> a positive natural atomic charge of  $0.631|e|$ . The covalent component of the Re–CO<sub>2</sub> bond is presented by the bonding  $\sigma(\text{Re}–\text{CO}_2)$  NBO which is constructed from the interaction of  $\text{sp}^{2.83}\text{d}^{2.52}$  hybrid orbitals (44.5% p and 39.7% d character) of Re with an  $\text{sp}^{1.62}$  hybrid (61.8% p-character) on the carbon donor atom of the CO<sub>2</sub> ligand and described as  $\sigma(\text{Re}–\text{O}) = 0.614h_{\text{Re}} + 0.789h_{\text{C}}$ . The occupancy or the  $\sigma(\text{Re}–\text{O})$  NBO is  $0.947|e|$ .

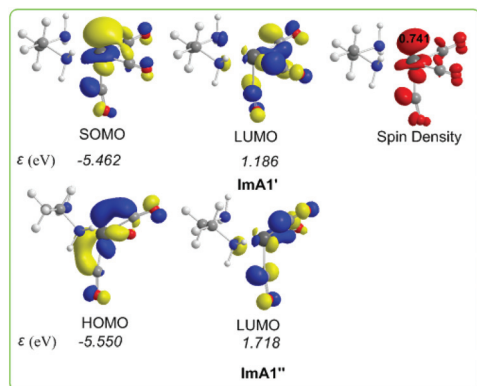
Successive protonations of the coordinated CO<sub>2</sub> yielding **ImC1''** and **ImD1''** are exergonic processes with estimated  $\Delta G$  values of  $-194.2$  and  $-154.4 \text{ kcal mol}^{-1}$  respectively. **ImD1''** releasing H<sub>2</sub>O in an exergonic process ( $\Delta G = -15.3 \text{ kcal mol}^{-1}$ ) affords the cationic [Re(en)(CO)<sub>4</sub>]<sup>+</sup> complex **ImE''** which upon 2e<sup>−</sup> reduction releases the desired CO product and regenerates the catalytically active **ImA1''** species.

In summary reaction steps B1' and B1'' are less favoured than reaction step B1 due to higher energy consumption to achieve the release of the desired CO product from the cationic [Re(en)(CO)<sub>4</sub>]<sup>+</sup> complex by 2e<sup>−</sup> reductions. Furthermore, the CO<sub>2</sub> insertion is more feasible than CO<sub>2</sub> capture illustrating the crucial role of the sacrificial electron donor TEOA in the photocatalytic reduction of CO<sub>2</sub> to CO catalyzed by the Re(en)(CO)<sub>3</sub>Cl complex.

## Reaction step B2

Reaction step B2 starts with the five-coordinated [(en)(CO)<sub>2</sub>ReCl]<sup>−</sup> intermediate that resulted from the dissociation of one of the equatorial CO ligands of the excited T<sub>1</sub> triplet state (step A2). It is important to note that Jean-Marie Lehn and co-workers<sup>19</sup> showed, by a set of experiments concerning the carbonyl ligand exchange in [*fac*-Re(bpy)(CO)<sub>3</sub>Cl], that reductive quenching of the excited state of the [Re(bpy)(CO)<sub>3</sub>Cl] complex by tertiary amines gives an unstable five-coordinated [Re(bpy)(CO)<sub>2</sub>Cl] species upon labilization of a CO ligand.

Two possible catalytic cycles following reaction steps B2' and B2'' have been thoroughly investigated. The reaction steps for the photocatalytic reduction of CO<sub>2</sub> to CO catalyzed by the five-coordinated [(en)(CO)<sub>2</sub>ReCl]<sup>−</sup>, **ImA2** intermediate are depicted schematically in Fig. 7.



**Scheme 3** 3D plots of the frontier molecular orbitals (FMOs) and spin density distribution of **ImA1'** and **ImA1''** catalytic species calculated at the wB97XD/Def2-TZVP level in the DMF solution.





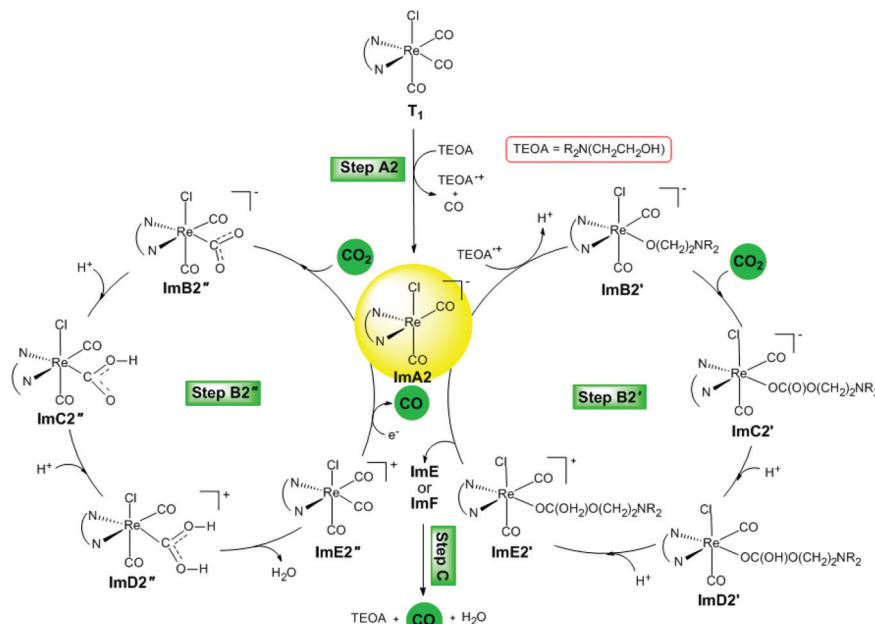
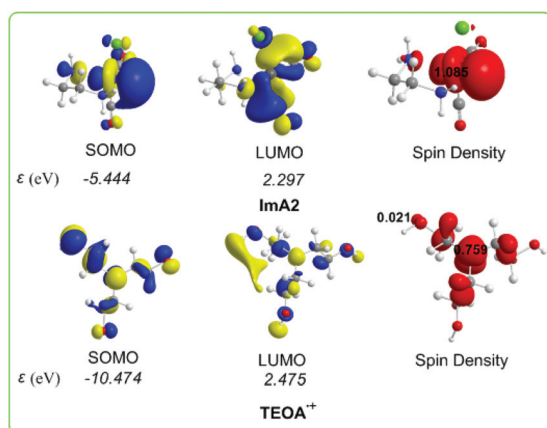


Fig. 7 Proposed catalytic cycles for steps B2' and B2'' calculated at the wB97XD/Def2-TZVP level in the DMF solution.



Scheme 4 3D plots of the frontier molecular orbitals (FMOs) and spin density distribution of **ImA2** and **TEOA<sup>+</sup>** species calculated at the wB97XD/Def2-TZVP level of theory.

Scheme 4 shows the 3D plots of the frontier molecular orbitals (FMOs) and spin density distribution of **ImA2** and **TEOA<sup>+</sup>** species.

The SOMO of **ImA2** is composed mainly of 84.8% Re  $\text{sp}_2\text{d}_{z^2}$  hybrid orbitals, while the LUMO is composed of 72.0% Re  $\text{pd}$  hybrid orbitals and 8.3% 2p AOs of the carbon atom of the CO ligand in *trans* position to the chloride ligand. A spin density of 1.085|e| is localized on the Re central atom. According to the NBO population analysis, the Re metal centre acquires a negative natural atomic charge of  $-0.552|e|$ . The SOMO of **TEOA<sup>+</sup>** is mainly localized on one of the  $\text{CH}_2\text{CH}_2\text{OH}$  alkyl groups (50.0%) mainly on the O atom (32.5%) and on the nitrogen atom (16.5%) of TEOA. Both the nature of the FMOs

and the negative natural atomic charge on the Re metal centre could not support covalent interactions with the  $\text{CO}_2$  ligand, thereby the capture of  $\text{CO}_2$  by **ImA2** might be supported mainly by non-covalent interactions. Indeed the interaction energy for the **ImA2**... $\text{CO}_2$  association is only 4.0 kcal mol<sup>-1</sup> calculated according to the wB97XD/Def2-TZVP/PCM computational protocol. However the nature of the FMOs of **ImA2** and **TEOA<sup>+</sup>** allows SOMO<sub>**ImA2**</sub>-SOMO<sub>**TEOA<sup>+</sup>**</sub> interactions giving rise to the covalent component of the **ImA2**-**TEOA** bonding upon the formation of a bonding  $\sigma(\text{Re}-\text{OCH}_2\text{CH}_2\text{NR}_2)$  NBO.

### Reaction step B2': $\text{CO}_2$ insertion

The geometric and free energy reaction profiles calculated for step B2' are shown in Fig. 8.

Perusal of Fig. 8 shows that an  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}^\bullet$  radical is coordinated to the Re metal centre of the catalytically active **ImA2** species which disposes of a vacant coordination site yielding **ImB2'**. The interaction of  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}^\bullet$  with **ImA2** is exergonic ( $\Delta G = -176.4$  kcal mol<sup>-1</sup>). The NBO population analysis of **ImB2'** revealed that the  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}^\bullet$  species is coordinated to the Re metal centre of **ImA2** forming a  $\text{Re}-\text{OCCH}_2\text{CH}_2\text{NR}_2$  bond with a WBI(Re-O) of 0.604. The covalent component of the Re-O bond is described by the bonding  $\sigma(\text{Re}-\text{O})$  NBO, with an occupancy of 1.929|e|. The  $\sigma(\text{Re}-\text{O})$  NBO results from the interaction of the  $\text{sp}^{2.89}\text{d}^{4.15}$  hybrid orbitals (35.8% p and 51.4% d character) of Re with an  $\text{sp}^{2.16}$  hybrid (68.3% p-character) of the O donor atom of the  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}^\bullet$  ligand and is described as  $\sigma(\text{Re}-\text{O}) = 0.399h_{\text{Re}} + 0.917h_{\text{O}}$ . The O atom of the coordinated  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}$  ligand acquires a negative natural atomic charge of  $-0.759|e|$  indicative of an anionic  $\text{R}_2\text{N}-\text{CH}_2\text{CH}_2\text{O}^-$  ligand. The Re metal center



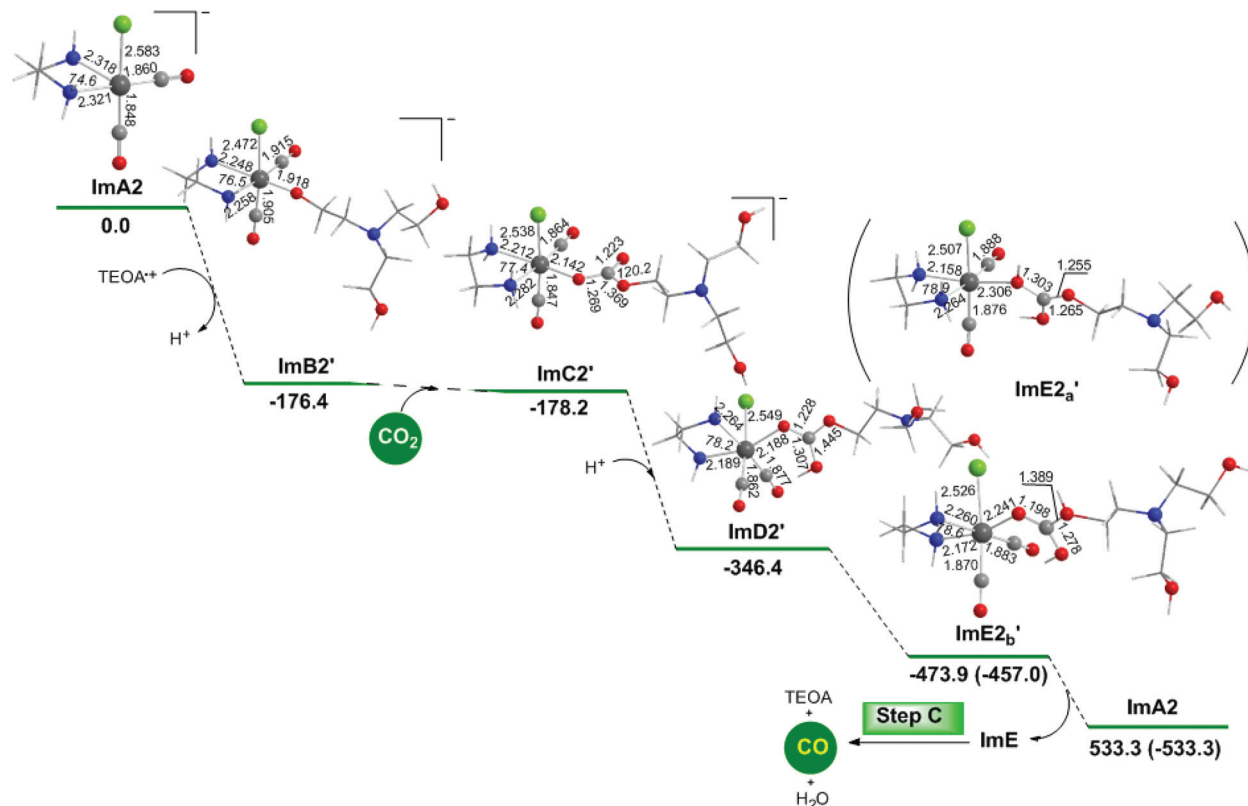
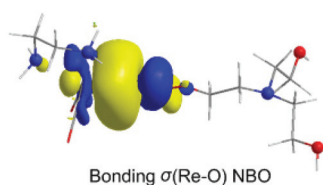


Fig. 8 Geometric and free energy,  $\Delta G$  (in  $\text{kcal mol}^{-1}$ ), reaction profiles for step B2' of the reduction of  $\text{CO}_2$  to CO catalyzed by the active  $[(\text{en})(\text{CO})_3\text{Re}]^-$  catalytic species, **ImA2**, calculated at the wB97XD/Def2-TZVP level in the DMF solution.



Scheme 5 3D plot of the bonding  $\sigma(\text{Re}–\text{O})$  NBO of **ImC2'** calculated at the wB97XD/Def2-TZVP level of theory.

in **ImB2'** acquires also a negative natural atomic charge of  $-0.249|e|$ .

Next insertion of a  $\text{CO}_2$  molecule into the  $\text{Re}–\text{OCH}_2\text{CH}_2\text{NR}_2$  transforms the  $\text{OCH}_2\text{CH}_2\text{NR}_2$  ligand to  $\text{OC}(\text{O})\text{OCH}_2\text{CH}_2\text{NR}_2$  yielding **ImC2'**. The insertion process is slightly exergonic ( $\Delta G = -1.8 \text{ kcal mol}^{-1}$ ). The  $\text{Re}–\text{OC}(\text{O})\text{OCH}_2\text{CH}_2\text{NR}_2$  coordination bond is a relatively weak  $\text{Re}–\text{O}$  dative bond with a  $\text{WBI}(\text{Re}–\text{O})$  of 0.485 described by the bonding  $\sigma(\text{Re}–\text{O})$  NBO as shown in Scheme 5 with an occupancy of  $1.934|e|$ .

The  $\sigma(\text{Re}–\text{O})$  NBO is constructed from the interaction of the  $\text{sp}^{3.00}\text{d}^{3.39}$  hybrid orbitals (40.5% p and 45.7% d character) of Re with an  $\text{sp}^{1.73}$  hybrid (63.3% p-character) of the O donor atom of the  $\text{R}_2\text{N}–\text{CH}_2\text{CH}_2\text{OC}(\text{O})\text{O}$  ligand and is described as  $\sigma(\text{Re}–\text{O}) = 0.364h_{\text{Re}} + 0.931h_{\text{O}}$ . The carbonate O atoms of the coordinated  $\text{R}_2\text{N}–\text{CH}_2\text{CH}_2\text{OC}(\text{O})\text{O}$  ligand acquiring negative

natural atomic charges of  $-0.744$ ,  $-0.666$  and  $-0.526|e|$  are susceptible to protonation, particularly the carboxylate O atoms yielding the mono-**ImD2'** and di-protonated **ImE2a'** or **ImE2b'** intermediates. **ImE2a'** and **ImE2b'** differ in the protonation sites of the coordinated  $\text{R}_2\text{N}–\text{CH}_2\text{CH}_2\text{OC}(\text{O})\text{O}$  ligand. The successive protonations are exergonic, and the estimated  $\Delta G$  values are  $-168.2 \text{ kcal mol}^{-1}$  and  $-127.5$  ( $-110.6$ )  $\text{kcal mol}^{-1}$ . Both **ImE2a'** and **ImE2b'** intermediates are unstable transient species directly releasing the cationic di-protonated  $[\text{R}_2\text{N}–\text{CH}_2\text{CH}_2\text{OC}(\text{OH})_2]^+$ , **ImE**, species and regenerating the catalytically active **ImA2** species. The **ImE** species participates in reaction step C (Fig. 1) that delivers the desired CO product.

### Reaction step B2': $\text{CO}_2$ capture

The geometric and free energy reaction profiles calculated for step B2' are shown in Fig. 9. The capture of  $\text{CO}_2$  by **ImA2** in reaction step B2' yielding **ImB2'** is slightly exergonic ( $-4.0 \text{ kcal mol}^{-1}$ ) and it is achieved by the nucleophilic attack of the electrophilic carbon atom of  $\text{CO}_2$  since the Re metal centre acquires a negative natural atomic charge of  $-0.507|e|$  and the C atom of the coordinated  $\text{CO}_2$  a positive natural atomic charge of  $0.737|e|$ . The NBO population analysis of **ImB2'** revealed that the  $\text{CO}_2$  molecule is coordinated to the Re metal centre of the **ImA2** catalytic species, forming a  $\text{Re}–\text{CO}_2$  bond with a  $\text{WBI}(\text{Re}–\text{CO}_2)$  of 0.634. The covalent component of the  $\text{Re}–\text{CO}_2$  bond is described by the bonding  $\sigma^a(\text{Re}–\text{CO}_2)$  and



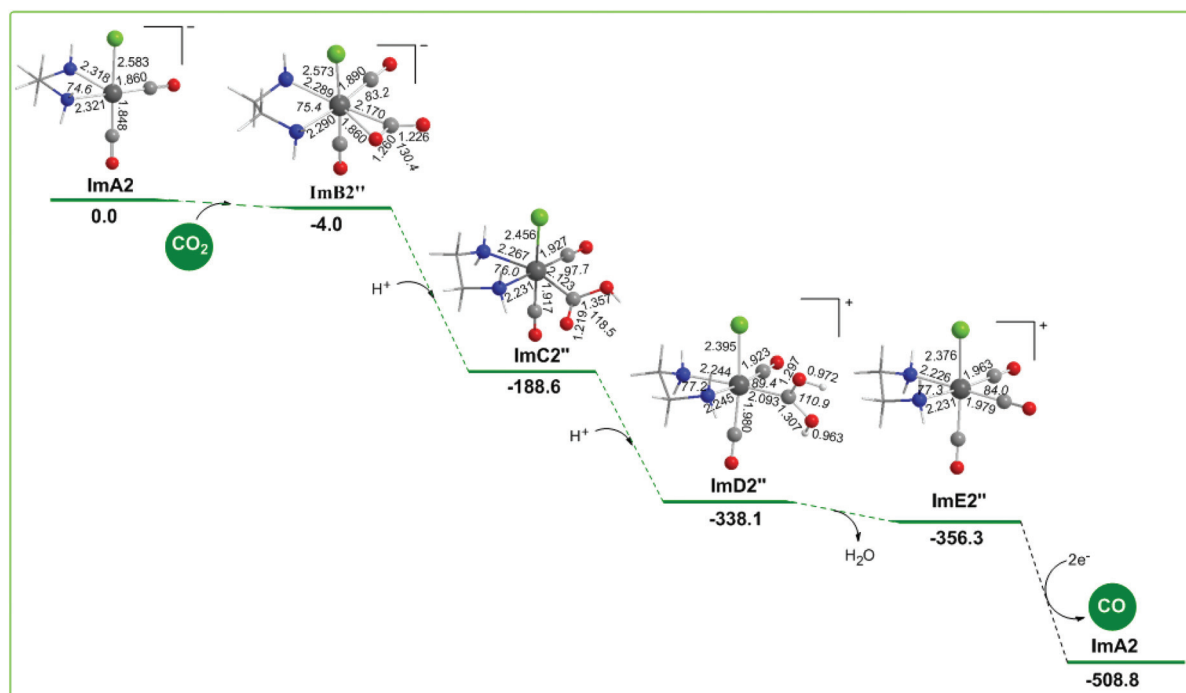


Fig. 9 Geometric and free energy,  $\Delta G$  (in  $\text{kcal mol}^{-1}$ ), reaction profiles for step B2'' of the reduction of  $\text{CO}_2$  to CO catalyzed by the active  $[(\text{en})(\text{CO})_3\text{Re}]^-$  catalytic species, **ImA2**, calculated at the wb97XD/Def2-TZVP level in the DMF solution.

$\sigma^{\beta}(\text{Re}-\text{CO}_2)$  NBOs. The  $\sigma^{\alpha}(\text{Re}-\text{CO}_2)$  NBO, with an occupancy of  $0.920|e|$ , results from the interaction of  $\text{sp}^{2.47}\text{d}^{2.27}$  hybrid orbitals (42.9% p and 39.4% d character) of Re with an  $\text{sp}^{1.79}$  hybrid (64.1% p-character) on the carbon donor atom of the  $\text{CO}_2$  ligand and is described as  $\sigma(\text{Re}-\text{CO}_2) = 0.610h_{\text{Re}} + 0.792h_{\text{C}}$ . The  $\sigma^{\beta}(\text{Re}-\text{CO}_2)$  NBO, with an occupancy of  $0.767|e|$ , results from the interaction of the  $\text{sp}^{4.25}\text{d}^{3.94}$  hybrid orbitals (46.2% p and 42.8% d character) of Re with an  $\text{sp}^{1.96}$  hybrid (66.2% p-character) on the carbon donor atom of the  $\text{CO}_2$  ligand and is described as  $\sigma^{\beta}(\text{Re}-\text{CO}_2) = 0.679h_{\text{Re}} + 0.735h_{\text{C}}$ . The O atoms of the coordinated  $\text{C}(\text{O})\text{O}$  ligand acquiring negative natural atomic charges of  $-0.646$  and  $-0.746|e|$  are susceptible to protonation. The first and second protonations transforming the coordinated  $\text{C}(\text{O})\text{O}$  ligand to  $\text{C}(\text{O})\text{OH}$  and  $\text{C}(\text{OH})\text{OH}$  ligands and yielding **ImC2''** and **ImD2''** intermediates respectively are strongly exergonic, with the estimated  $\Delta G$  values being  $-184.6$  and  $-149.5 \text{ kcal mol}^{-1}$ . **ImD2''** is a transient species directly decomposed to give  $\text{H}_2\text{O}$  and **ImE2''** formulated as the  $[\text{Re}(\text{en})(\text{CO})_3\text{Cl}]^+$  complex which by  $2e^-$  reduction releases the desired CO product and regenerates the starting **ImA2** complex. The two electrons are provided by the TEOA sacrificial electron donor. Note that the proposed catalytic cycle for step B2'' is exactly the same as a possible catalytic cycle for the photocatalytic reduction of  $\text{CO}_2$  to CO catalyzed by the  $[\text{Re}(\text{bpy})(\text{CO})_2\text{Cl}]$  catalytic species generated from the loss of a CO ligand from the reduced form of the  $[\text{Re}(\text{bpy})(\text{CO})_3\text{Cl}]$  complex proposed by Lehn and co-workers.<sup>19</sup>

In summary the free energy reaction profiles for reaction steps B2' and B2'' suggest that both the reaction pathways

would proceed in parallel, but the  $\text{CO}_2$  insertion pathway should be most favorable than the  $\text{CO}_2$  capture pathway, since in the latter the  $\text{CO}_2$  capture by the **ImA2** catalytic species is difficult and the release of CO from the cationic  $[\text{Re}(\text{en})(\text{CO})_3\text{Cl}]^+$ , **ImE2''** intermediate should be subject to  $2e^-$  reduction, that is the energy consuming process. The  $\text{CO}_2$  capture pathway is expected to be the most favorable reaction pathway in the electrochemical reduction of  $\text{CO}_2$  to CO mediated by the  $[\text{Re}(\text{en})(\text{CO})_3\text{Cl}]$  catalyst in line with the catalytic cycle for the reduction of  $\text{CO}_2$  to CO mediated by the  $[\text{Re}(\text{bpy})(\text{CO})_3\text{Cl}]$  catalyst proposed by Lehn and co-workers.<sup>19</sup>

## Conclusions

This study reveals the crucial role of the TEOA sacrificial electron donor in the photocatalytic reduction of  $\text{CO}_2 \rightarrow \text{CO}$  catalyzed by the  $[\text{Re}(\text{en})(\text{CO})_3\text{Cl}]$  complex. Despite its electron and proton donor abilities, TEOA can capture  $\text{CO}_2$  upon coordination to the Re metal centre of the "real" catalytic species  $[\text{Re}(\text{en})(\text{CO})_3]$ ,  $[\text{Re}(\text{en})(\text{CO})_3]^-$  and/or  $[\text{Re}(\text{en})(\text{CO})_2\text{Cl}]^-$  generated from the  $\text{T}_1$  state upon single and double reductive quenching by TEOA or photodissociation of a CO ligand. The oxidized  $\text{R}_2\text{NCH}_2\text{CH}_2\text{O}^\bullet$  radical of TEOA upon coordination with the Re metal centre of the catalytic species promotes the  $\text{CO}_2$  insertion in the  $\text{Re}-\text{OCH}_2\text{CH}_2\text{NR}_2$  bond forming stable intermediates that upon successive protonations yield unstable transient species that release CO and regenerate the catalytic species. In particular, in the  $\text{CO}_2$  insertion pathways, CO is delivered by



an unstable diprotonated  $[R_2NCH_2CH_2OC(OH)(OH)]^+$  species regenerating TEOA and the catalyst. The  $CO_2$  insertion reaction pathway is the favourable pathway for the photocatalytic reduction of  $CO_2 \rightarrow CO$  catalysed by the  $[Re(en)(CO)_3Cl]$  complex in the presence of TEOA manifesting its crucial role as an electron and proton donor, capturing  $CO_2$  and releasing CO. The  $CO_2$  capture reaction pathways involving direct capture of  $CO_2$  by the catalytic species is less favorable because the direct capture of  $CO_2$  is a difficult task and the release of CO from the  $[Re(en)(CO)_4]^+$  and  $[Re(en)(CO)_2Cl]^+$  intermediates formed upon successive protonation of the captured  $CO_2$  ligand corresponds to energy consuming processes. This is why the  $CO_2$  capture reaction pathways are favourable pathways in the electrochemical reduction of  $CO_2$  to CO mediated by the  $[Re(en)(CO)_3Cl]$  catalyst in line with the catalytic cycle proposed by Lehn and co-workers.<sup>19</sup> We believe that the present in-depth mechanistic investigations would obviously inspire the chemists towards new and more efficient, selective and stable catalytic systems for the  $CO_2$  insertion step and easily releasing the reduction product of CO.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The work has been performed under the Project HPC-EUROPA3 (INFRAIA-2016-1-730897), with the support of the EC Research Innovation Action under the H2020 Programme; in particular, the author gratefully acknowledges the support of Prof. M. Bickelhaupt, Department of Chemistry, Vrije Universiteit Amsterdam and the computer resources and technical support provided by the SURFsara supercomputing centre.

## Notes and references

- M. Aresta and D. Dibenedetto, *Dalton Trans.*, 2007, 2975–2992.
- M. Cokoja, C. Bruckmeier, B. Rieger, F. Kühnand and W. A. Hermann, *Angew. Chem., Int. Ed.*, 2011, **50**, 8510–8537.
- A. J. Morris, G. J. Meyer and E. Fujita, *Acc. Chem. Res.*, 2009, **42**, 1983–1994.
- J. A. Keith, K. A. Grice, C. P. Kubiak and E. A. Carter, *J. Am. Chem. Soc.*, 2013, **135**, 15823–15829.
- C. Riplinger, M. D. Sampson, A. M. Ritzmann, C. P. Kubiak and E. A. Carter, *J. Am. Chem. Soc.*, 2014, **136**, 16285–16298.
- K. S. Rawat, A. Mahata, I. Choudhuri and B. Pathak, *J. Phys. Chem. C*, 2016, **120**(16), 8821–8831.
- D. H. Apaydin, S. Schlager, E. Portenkirchner, S. Niyazi and N. S. Sariciftci, *ChemPhysChem*, 2017, **18**, 3094–3116.
- N. Elgrishi, M. B. Chambers, X. Wang and M. Fontecave, *Chem. Soc. Rev.*, 2017, **46**, 761–796.
- K. E. Dalle, J. Warnan, J. J. Leung, B. Reuillard, I. S. Karmel and E. Reisner, *Chem. Rev.*, 2019, **119**, 2752–2875.
- Y. Yamazaki, H. Takeda and O. Ishitani, *J. Photochem. Photobiol., C*, 2015, **25**, 106–137.
- Y. Kuramochi, O. Ishitani and H. Ishida, *Coord. Chem. Rev.*, 2018, **373**, 333–356.
- H. Koizumi, H. Chiba, A. Sugihara, M. Iwamura, K. Nozaki and O. Ishitani, *Chem. Sci.*, 2019, **10**, 3080–3088.
- Y. Asai, H. Katsuragi, K. Kita, T. Tsubomura and Y. Yamazaki, *Dalton Trans.*, 2020, **49**, 4277–4292.
- I. Choudhuri and D. G. Truhlar, *J. Phys. Chem. C*, 2020, **124**(16), 8504–8513.
- S. Gong, J. Fan, V. Cecenc, C. Huang, Y. Min, Q. Xu and H. Li, *Chem. Eng. J.*, 2021, **405**, 126913–112627.
- H. Yuan, B. Cheng, J. Lei, L. Jiang and Z. Han, *Nat. Commun.*, 2021, **12**, 1835–1844.
- J.-M. Lehn and R. Ziessel, *Proc. Natl. Acad. Sci. U. S. A.*, 1982, **79**, 701–704.
- J. Hawecker, J.-M. Lehn and R. Ziessel, *J. Chem. Soc., Chem. Commun.*, 1983, 536–538.
- J. Hawecker, J.-M. Lehn and R. Ziessel, *Helv. Chim. Acta*, 1986, **69**, 1990–2012.
- C. Kütal, M. A. Weber, G. Ferraudi and D. A. Geiger, *Organometallics*, 1985, **4**, 2161–2166.
- C. Kütal, M. A. Weber, G. Ferraudi and D. A. Geiger, *Organometallics*, 1987, **6**, 553–557.
- J. M. Smieja and C. P. Kubiak, *Inorg. Chem.*, 2010, **49**, 9283.
- J. M. Smieja, E. E. Benson, B. Kumara, K. A. Grice, C. S. Seua, J. M. Miller, J. M. Mayer, P. Clifford and C. P. Kubiak, *Proc. Natl. Acad. Sci. U. S. A.*, 2012, **109**, 15646–15650.
- K. A. Grice, N. X. Gu, M. D. Sampson and C. P. Kubiak, *Dalton Trans.*, 2013, **42**, 8498–8503.
- E. E. Benson, K. A. Grice, J. M. Smieja, P. Clifford and C. P. Kubiak, *Polyhedron*, 2013, **58**, 229–234.
- J. Agarwal, E. Fujita, H. F. Schaefer III and J. T. Muckerman, *J. Am. Chem. Soc.*, 2012, **134**, 5180–5186.
- H. Takeda, K. Koike, H. Inoue and O. Ishitani, *J. Am. Chem. Soc.*, 2008, **130**, 2023–2031.
- T. Morimoto, T. Nakajima, S. Sawa, R. Nakanishi, D. Imori and O. Ishitani, *J. Am. Chem. Soc.*, 2013, **135**, 16825.
- P. Kurz, B. Probst, B. Spingler and R. Alberto, *Eur. J. Inorg. Chem.*, 2006, 2966–2974.
- Y. Kuramochi, K. Fukaya, M. Yoshida and H. Ishida, *Chem. – Eur. J.*, 2015, **21**, 10049–10060.
- M. J. Frisch, *et al.*, *Gaussian 16, Revision C.01*, Gaussian, Inc., Wallingford, CT, 2010.
- S. Grimme, *J. Comput. Chem.*, 2006, **27**, 1787–1799.
- D. Becke, *J. Chem. Phys.*, 1997, **107**, 8554–8560.
- D. Chai and M. Head-Gordon, *Phys. Chem. Chem. Phys.*, 2008, **10**, 6615–6620.
- Q. Wu and W. T. Yang, *J. Chem. Phys.*, 2002, **116**, 515–524.
- Y. Minenkov, A. Singstad, O. Occhipinti and V. Jensen, *Dalton Trans.*, 2012, **41**, 5526–5541.





- 37 J. Tomasi, B. Mennucci and R. Cammi, *Chem. Rev.*, 2005, **105**, 2999–3093.
- 38 A. E. Reed, L. A. Curtiss and F. Weinhold, *Chem. Rev.*, 1988, **88**, 899–926.
- 39 F. Weinhold, in *The Encyclopedia of Computational Chemistry*, ed. P. V. R. Schleyer, John Wiley & Sons, Chichester, U.K., 1998.
- 40 S. J. A. van Gisbergen, F. Kootstra, P. R. T. Schipper, O. V. Gritsenko, J. G. Snijders and E. J. J. Baerends, *Phys. Rev. A*, 1998, **57**, 2556–2571.
- 41 C. Jamorski, M. E. Casida and D. R. Salahub, *J. Chem. Phys.*, 1996, **104**, 5134–5147.
- 42 R. Bauernschmitt and R. Ahlrichs, *Chem. Phys. Lett.*, 1996, **256**, 454–464.

