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# Promoting photocatalytic CO<sub>2</sub> reduction through facile electronic modification of N-annulated perylene diimide rhenium bipyridine dyads†

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The development of CO<sub>2</sub> conversion catalysts has become paramount in the effort to close the carbon loop. Herein, we report the synthesis, characterization, and photocatalytic CO<sub>2</sub> reduction performance for a series of N-annulated perylene diimide (NPDI) tethered Re(bpy) supramolecular dyads [Re(bpy-C2-NPDI-R)], where R = -H, -Br, -CN, -NO<sub>2</sub>, -OPh, -NH<sub>2</sub>, or pyrrolidine (-NR<sub>2</sub>). The optoelectronic properties of these Re(bpy-C2-NPDI-R) dyads were heavily influenced by the nature of the R-group, resulting in significant differences in photocatalytic CO<sub>2</sub> reduction performance. Although some R-groups (*i.e.* -Br and -OPh) did not influence the performance of CO<sub>2</sub> photocatalysis (relative to -H; TON<sub>CO</sub> ~60), the use of an electron-withdrawing -CN was found to completely deactivate the catalyst (TON<sub>CO</sub> < 1) while the use of an electron-donating -NH<sub>2</sub> improved CO<sub>2</sub> photocatalysis four-fold (TON<sub>CO</sub> = 234). Despite being the strongest EWG, the -NO<sub>2</sub> derivative exhibited good photocatalytic CO<sub>2</sub> reduction abilities (TON<sub>CO</sub> = 137). Using a combination of CV and UV-vis-nIR SEC, it was elucidated that the -NO<sub>2</sub> derivative undergoes an *in situ* transformation to -NH<sub>2</sub> under reducing conditions, thereby generating a more active catalyst that would account for the unexpected activity. A photocatalytic CO<sub>2</sub> mechanism was proposed for these Re(bpy-C2-NPDI-R) dyads (based on molecular orbital descriptions), where it is rationalized that the photoexcitation pathway, as well as the electronic driving-force for NPDI<sup>2-</sup> to Re(bpy) electron-transfer both significantly influence photocatalytic CO<sub>2</sub> reduction. These results help provide rational design principles for the future development of related supramolecular dyads.

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## Introduction

The adverse effects on climate change related to increased anthropogenic CO<sub>2</sub> emissions has inspired the utilization of excess CO<sub>2</sub> as a sustainable feedstock for value-added chemicals and fuels.<sup>1,2</sup> While the activation of CO<sub>2</sub> is kinetically unfavorable, it can be readily accomplished electro-/photocatalytically *via* proton-couple multielectron chemical reductions.<sup>3,4</sup> Consequently, the development of capable molecular electro-/photocatalysts has mainly focused on improving the efficiency and selectivity of the CO<sub>2</sub> conversion process.<sup>5-9</sup> Among the many comprehensively studied molecular catalyst systems, Re(2,2'-bipyridine)(CO)<sub>3</sub>Cl [Re(bpy)] is notable for its highly selective CO<sub>2</sub>-to-CO conversion.<sup>10</sup> The versatile bpy ligand has been modified with a variety of substituents to change both the electronic properties and/or the second-sphere H-bonding character of the catalyst.<sup>11-18</sup> And while Re(bpy) alone can be used as an effective CO<sub>2</sub> reduction photocatalyst,<sup>19-24</sup> the

photocatalytic CO<sub>2</sub> reduction performance is greatly enhanced *via* the direct functionalization of Re(bpy) with photosensitizing (PS) units.<sup>25-31</sup>

The development of ruthenium(II) diimine photosensitized Re(bpy) supramolecular dyads has been extensively reported by the Ishitani group.<sup>5,25-31</sup> These Ru<sup>II</sup>-Re<sup>I</sup> dyads make use of a Z-scheme architecture whereby the photoexcited electrons of the Ru<sup>II</sup>-moiety are reductively quenched and subsequently transferred to the Re<sup>I</sup> catalyst center to enable CO<sub>2</sub> reduction. To facilitate efficient electron-transfer (eT) and CO<sub>2</sub> photocatalysis, several supra-molecular dyad design principles have been established. First, the photoexcited electron should be localized near the tethering portion between the PS and the catalyst.<sup>5</sup> Second, the tether between the PS and catalyst moieties should be as short as possible (without being through-conjugated) to enable rapid intramolecular eT.<sup>27-30</sup> Third, increasing the molar absorptivity of the PS-moiety (*i.e.* by incorporating multiple PS units) can improve the quantum efficiency capabilities and the ensuing eT dynamics of the supramolecular dyad.<sup>31</sup> Restricted by the first two design principles, attempts to improve the quantum efficiency of these Re(bpy) dyads has been made by using more strongly absorbing PS units, such as porphyrins,<sup>32-37</sup> naphthalimide,<sup>38,39</sup> naphthalene diimide,<sup>40-42</sup> and perylene diimide (PDI).<sup>43,44</sup> Although the photophysical dynamics of these

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dyads appear fundamentally well-understood, only a handful have been properly evaluated as CO<sub>2</sub> reduction photocatalysts (see ESI, Table S1†).<sup>35–39</sup>

Recently, we reported on four N-annulated perylene diimide (NPDI) functionalized Re(bpy) dyads as CO<sub>2</sub> reduction electrocatalysts.<sup>45</sup> Our investigation of these Re(bpy)–NPDI dyads revealed that the PS unit (NPDI) functions as an electron-reservoir for Re(bpy), enabling efficient CO<sub>2</sub> reduction at an overpotential 300 mV lower than conventional Re(bpy)-type electrocatalysts. Moreover, it was also elucidated that the tether length between Re(bpy) and NPDI governs which CO<sub>2</sub> reduction mechanism is preferred for the supramolecular dyad(s), where the ethyl-linked Re(bpy)–NPDI dyad possessed the greatest degree of electronic communication. These promising results from our initial Re(bpy)–NPDI dyads led us to hypothesize that eT from the electron-reservoir to the Re(bpy) catalyst could be improved by electronically modifying NPDI in two different ways. It was theorized that the introduction of electron withdrawing groups (EWGs) on NPDI may inductively stabilize the entire dyad, thus enabling more efficient eT by increasing the overall electron affinity of Re(bpy)-moiety. Alternatively, the use of electron donating groups (EDGs) on NPDI could make the electron-reservoir more electron-rich and thus more willing to transfer electrons to the Re(bpy)-moiety. To determine which of these two opposing hypotheses was correct, a series of electronically modified ethyl-linked Re(bpy)–NPDI dyads [Re(bpy-C2-NPDI-R)] (where R = –H, –Br, –CN, –NO<sub>2</sub>, –OPh, –NH<sub>2</sub>, or –NR<sub>2</sub>) were designed (Fig. 1, left) and, for the first time, their photocatalytic CO<sub>2</sub>-to-CO reduction performance was evaluated. It was revealed that installing an EDG, such as –NH<sub>2</sub>, led to a four-fold enhancement in turnover numbers of CO (TON<sub>CO</sub>), with respect to the benchmark Re(bpy-C2-NPDI-H) dyad. A mechanism based on molecular orbital (MO) energy levels is proposed to explain the observed differences in photocatalytic CO<sub>2</sub> reduction

performance for these dyads caused by the installation of EWGs and EDGs on NPDI.

## Results & discussion

### Synthesis & characterization

The synthesis of all azide-ethyl-NPDI precursors (N<sub>3</sub>-C2-NPDI-R) starts from the HNPDI synthon (Scheme 1).<sup>47–49</sup> The pyrrolic nitrogen was alkylated with 1,2-dibromoethane and a terminal azide was installed using S<sub>N</sub>2 chemistry to give N<sub>3</sub>-C2-NPDI-H.<sup>45,47</sup> Treatment of N<sub>3</sub>-C2-NPDI-H with either fuming HNO<sub>3</sub> (at –78 °C) or Br<sub>2</sub> (at 20 °C) affords the N<sub>3</sub>-C2-NPDI-NO<sub>2</sub> and N<sub>3</sub>-C2-NPDI-Br precursors, respectively. When refluxed with pyrrolidine, the N<sub>3</sub>-C2-NPDI-Br precursor can be converted to N<sub>3</sub>-C2-NPDI-NR<sub>2</sub>. Alternatively, reacting N<sub>3</sub>-C2-NPDI-Br with an excess of phenol and K<sub>2</sub>CO<sub>3</sub> in *N,N*-dimethylformamide (DMF; at 80 °C) generates N<sub>3</sub>-C2-NPDI-OPh. Attempts to cyano-functionalize N<sub>3</sub>-C2-NPDI using the Rosenmund-von Braun reaction conditions resulted in the elimination of the terminal azide. Consequently, N<sub>3</sub>-C2-NPDI-CN had to be synthesized starting from HNPDI. After selectively brominating HNPDI,<sup>50</sup> HNPDI-Br was reacted with excess CuCN in refluxing DMF to yield HNPDI-CN. This intermediate was then alkylated and azide-functionalized to give the N<sub>3</sub>-C2-NPDI-CN precursor.

Next, these six N<sub>3</sub>-C2-NPDI-R precursors were linked to Re(5-ethynyl-2,2'-bipyridine)(CO)<sub>3</sub>Cl [Re(ethynyl-bpy)] using a standard copper catalyzed azide-alkyne cycloaddition (CuAAC) procedure (see ESI Section II for more details†).<sup>45,47</sup> While most Re(bpy-C2-NPDI-R) dyads (where R = –H, –Br, –CN, –NR<sub>2</sub>, and –OPh) were obtained in excellent yields as the exclusive product, the CuAAC reaction between N<sub>3</sub>-C2-NPDI-NO<sub>2</sub> and Re(ethynyl-bpy) generated both Re(bpy-C2-NPDI-NO<sub>2</sub>) (major product) and Re(bpy-C2-NPDI-NH<sub>2</sub>) (minor product). To obtain each catalyst selectively, two additional CuAAC protocols were

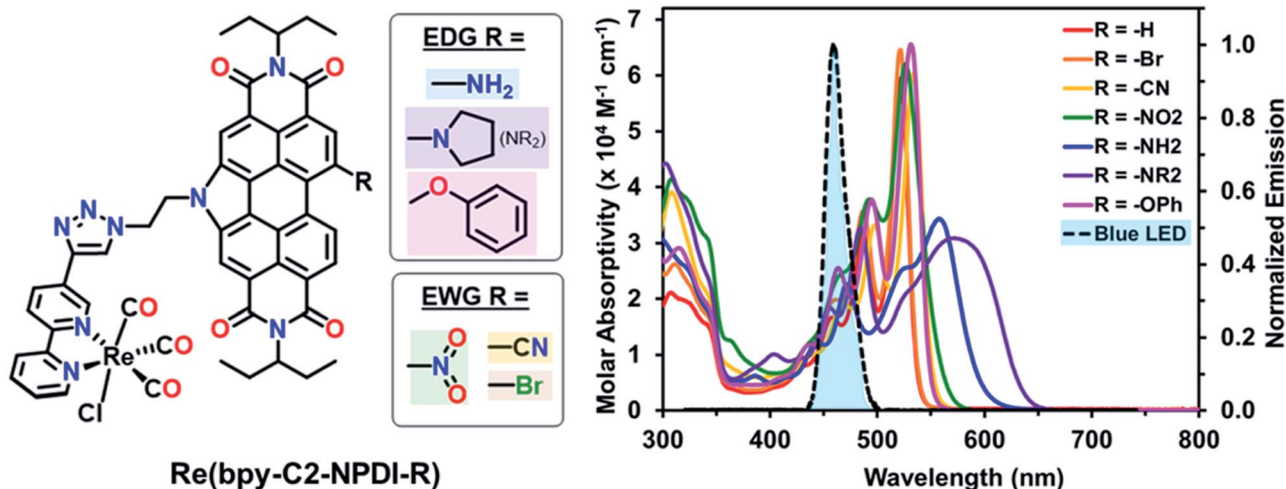
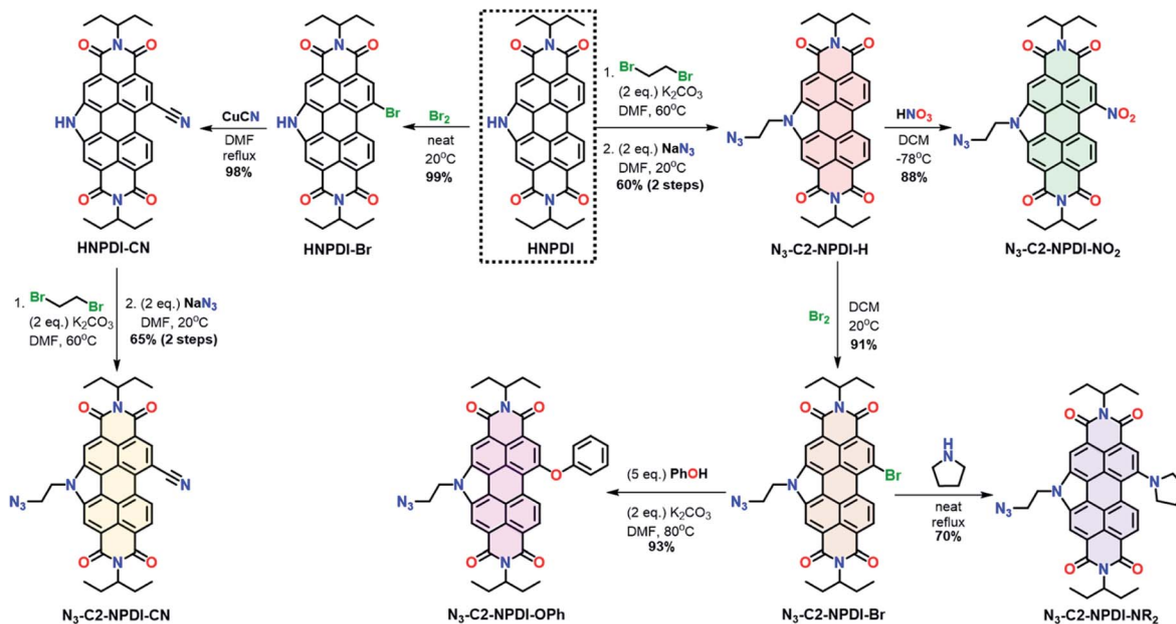


Fig. 1 (Left) chemical structures and (right) UV-vis absorption spectra of Re(bpy-C2-NPDI-R) dyads in CHCl<sub>3</sub> (~10<sup>–5</sup> M), where R = –H (red), R = –Br (orange), R = –CN (yellow), R = –NO<sub>2</sub> (green), R = –NH<sub>2</sub> (blue), R = –NR<sub>2</sub> (purple), and R = –OPh (magenta). The optical profiles of all Re(bpy-C2-NPDI-R) dyads overlap with the emission spectrum of the blue LED ( $\lambda = 470 \pm 30$  nm; 4 mW cm<sup>–2</sup>) used for photocatalytic CO<sub>2</sub> reduction testing.





Scheme 1 Synthesis of  $N_3$ -C2-NPDI-R precursors that are converted into Re(bpy-C2-NPDI-R) dyads.

developed. The selective synthesis of **Re(bpy-C2-NPDI-NO<sub>2</sub>)** was accomplished by changing the catalyst from CuSO<sub>4</sub> (with sodium ascorbate) to CuI, suggesting that the sub-stoichiometric sodium ascorbate used under our standard CuAAC conditions acted as a reducing agent.<sup>51–53</sup> With this insight, **Re(bpy-C2-NPDI-NH<sub>2</sub>)** was afforded exclusively using catalytic CuSO<sub>4</sub> and a stoichiometric excess of sodium ascorbate. The identity of all **Re(bpy-C2-NPDI-R)** dyads was confirmed using <sup>1</sup>H and <sup>13</sup>C NMR spectroscopies, as well as MALDI-TOF mass spectrometry and CHN elemental analysis (Fig. S1–S54<sup>†</sup>). Note, each specific **Re(bpy-C2-NPDI-R)** dyad will henceforth be referred to by their R-group only (where R = –H, –Br, –CN, –NO<sub>2</sub>, –NH<sub>2</sub>, –NR<sub>2</sub>, and –OPh).

The optical properties of these **Re(bpy-C2-NPDI-R)** dyads were probed using both UV-visible-near-infrared (UV-vis-nIR) absorption and Fourier transform infrared (FTIR) spectroscopies. Visually, the optical properties of all **Re(bpy-C2-NPDI-R)** dyads were dominated by the NPDI-moiety (Fig. 1 and S55–S56<sup>†</sup>). Relative to –H ( $\lambda_{\max} = 521$ ), the installation of –Br, –CN, –NO<sub>2</sub>, and –OPh groups on NPDI all caused a minor bathochromic shift of the absorption profile ( $\Delta\lambda_{\max} = 1–13$  nm) while maintaining most of the vibronic transitions of the parent complex. As for the –NH<sub>2</sub> and –NR<sub>2</sub> dyads, the optical spectra were significantly red-shifted ( $\lambda_{\max} = 558$  and 572 nm, respectively) and the molar absorptivity also decreased (from  $\sim 60\,000$  to  $\sim 40\,000$  M<sup>–1</sup> cm<sup>–1</sup>). The broadened absorption of –NH<sub>2</sub> and –NR<sub>2</sub> near the  $\lambda_{\max}$  is consistent with the donation of electron-density from the lone-pair of the amino nitrogen into the NPDI  $\pi$ -system.<sup>54–56</sup> Notably, all dyads exhibit an absorption peak around  $\lambda = 341–343$  nm, characteristic of a Re(bpy)-based metal-to-ligand charge transfer band.<sup>57</sup> By FTIR, all **Re(bpy-C2-NPDI-R)** dyads possessed three carbonyl stretching frequencies ( $\nu_{\text{co}} \sim 1900, 1925, \text{ and } 2025$  cm<sup>–1</sup>), which is consistent with

other Re(bpy) catalysts (Fig. S57<sup>†</sup>).<sup>11–18,58</sup> Together, these data confirm successful **Re(bpy-C2-NPDI-R)** dyad formation.

The electrochemical properties of these **Re(bpy-C2-NPDI-R)** dyads were next evaluated using cyclic voltammetry (CV). CV analysis was first performed in CH<sub>2</sub>Cl<sub>2</sub> (Fig. 2A and S65<sup>†</sup>), with all reported redox events being referenced to the Fe<sup>+/0</sup> internal standard. Under an atmosphere of argon, all **Re(bpy-C2-NPDI-R)** dyads exhibited four reduction and two oxidation redox processes. The first two reversible reductions may be assigned to the NPDI<sup>–/0</sup> and NPDI<sup>2–/1–</sup> redox couples while the third and fourth reductions correspond to the quasi-reversible bpy<sup>•–/0</sup> and the irreversible Re<sup>0/I</sup> redox events, respectively.<sup>11–18,45</sup> With respect to the oxidation events, the irreversible Re<sup>II/I</sup> redox process remains consistently near  $E_p \approx +1.0$  V. The quasi-reversible NPDI-based oxidation event, on the other hand, underwent dramatic shifts in potential depending on the nature of the electronic substituent.

Based on the presented CV data (Fig. 2A, S64 and S65<sup>†</sup>), it was observed that the installation of EWGs (*i.e.* –Br, –CN, and –NO<sub>2</sub>) on NPDI caused both the redox events assigned to the lowest unoccupied molecular orbital (LUMO) and the highest occupied molecular orbital (HOMO) energy levels of NPDI to shift to more positive potentials. In all cases, the electronic bandgap was narrowed because the LUMO energy level was more significantly perturbed than the HOMO energy level. Conversely, the installation of EDGs (*i.e.* –OPh, –NH<sub>2</sub>, and –NR<sub>2</sub>) on NPDI caused the redox events associated with NPDI HOMO and LUMO energy levels to shift to more negative potentials. Once again, the electronic bandgap was decreased mainly due to the more substantial effects experienced by the HOMO energy level. The observed shifts in  $E_{1/2}$  for first NPDI-R based reduction and oxidation events (Fig. 2B and C), relative to –H, show good Hammett parameter correlation.<sup>46</sup> Similar



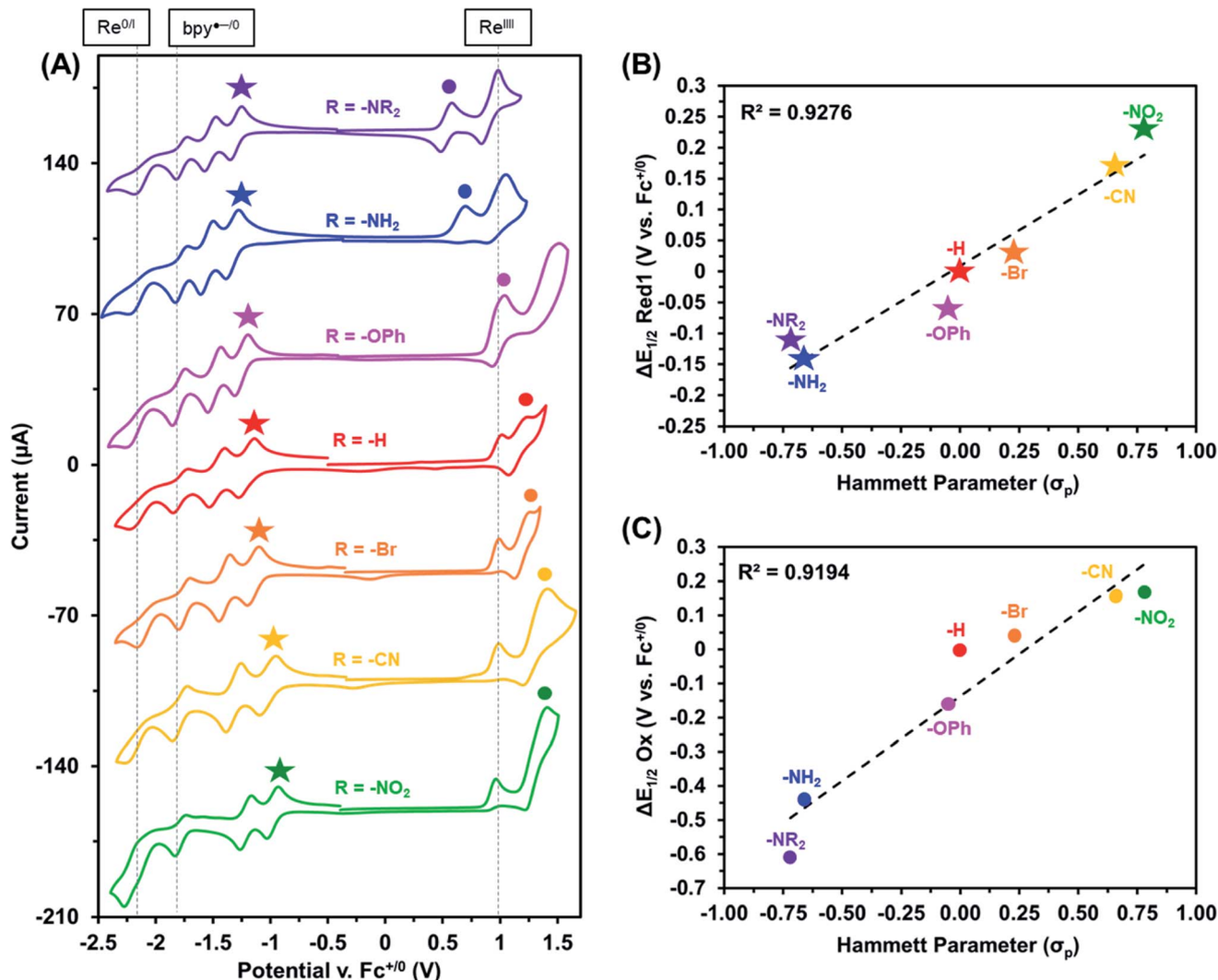


Fig. 2 Cyclic voltammograms (A) of  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads where  $R = -\text{NR}_2$  (purple),  $-\text{NH}_2$  (blue),  $-\text{OPh}$  (magenta),  $-\text{H}$  (red),  $-\text{Br}$  (orange),  $-\text{CN}$  (yellow), and  $-\text{NO}_2$  (green). All measurements were recorded at  $100 \text{ mV s}^{-1}$ , under argon in  $\text{CH}_2\text{Cl}_2$  with  $0.1 \text{ M TBAPF}_6$  supporting electrolyte (WE = glassy carbon, CE = Pt-wire, RE = Ag/AgCl, and  $\text{Fc}^{+/0}$  as internal reference standard). Correlation diagrams plot the observed shifts in  $E_{1/2}$  (relative to  $R = -\text{H}$ ) for the first reduction (B) and oxidation (C) as a function of R-group Hammett parameter.<sup>46</sup>

correlations were previously reported for electronically-substituted  $\text{Re}(4,4'\text{-R-bpy})$  complexes by Kubiak *et al.*,<sup>11</sup> where EWGs shifted redox events more positively and EDGs shifted redox events more negatively. In principle, combining these two relationships could assist with proper energy level matching when designing future supramolecular dyads based on the  $\text{Re}(\text{bpy-C2-NPDI-R})$  architecture.

Next, the electrochemistry of these  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads were assessed in DMF, where only the reduction processes were measured due to solvent window effects.<sup>59</sup> Under argon, all  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads exhibited the same four previously assigned reduction events (*vide supra*), where solvent effects caused the potential of most redox events to shift positively by  $\sim 50\text{--}100 \text{ mV}$ , relative to  $\text{CH}_2\text{Cl}_2$  (Fig. S66†).<sup>60</sup> Upon measuring CVs at variable scan rates, each dyad displayed a diffusion-limited current response when fitted to the Randles-Sevcik equation (see ESI, eqn (i)†), with a calculated diffusion

coefficient between  $D = 2.3\text{--}4.4 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$  (Fig. S69–S74†). Upon subjecting these  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads to an atmosphere of  $\text{CO}_2$  (in DMF), a moderate CV current enhancement was observed underneath the fourth redox couple (Fig. 3, red traces with  $E_{\text{cat}/2} \approx -2.1 \text{ V}$ ). When a catalyst is supplied with the appropriate combination of substrates at a sufficient applied potential, a CV current enhancement is often observed as a result of initiating an electron-consuming catalytic process.<sup>61,62</sup> Moreover, these CV current enhancements may be monitored as a function of catalyst or proton-source concentration to gather preliminary mechanistic data on the catalytic process (see ESI, eqn (ii)†).<sup>63,64</sup>

When the  $\text{Re}(\text{bpy-C2-NPDI-R})$  concentration was varied, a linear CV current enhancement response was also observed for most dyads (Fig. S68†). Similarly, when proton-source 2,2,2-trifluoroethanol (TFE) was incrementally added to a  $\text{CO}_2$ -saturated  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyad solution, it induced a CV current



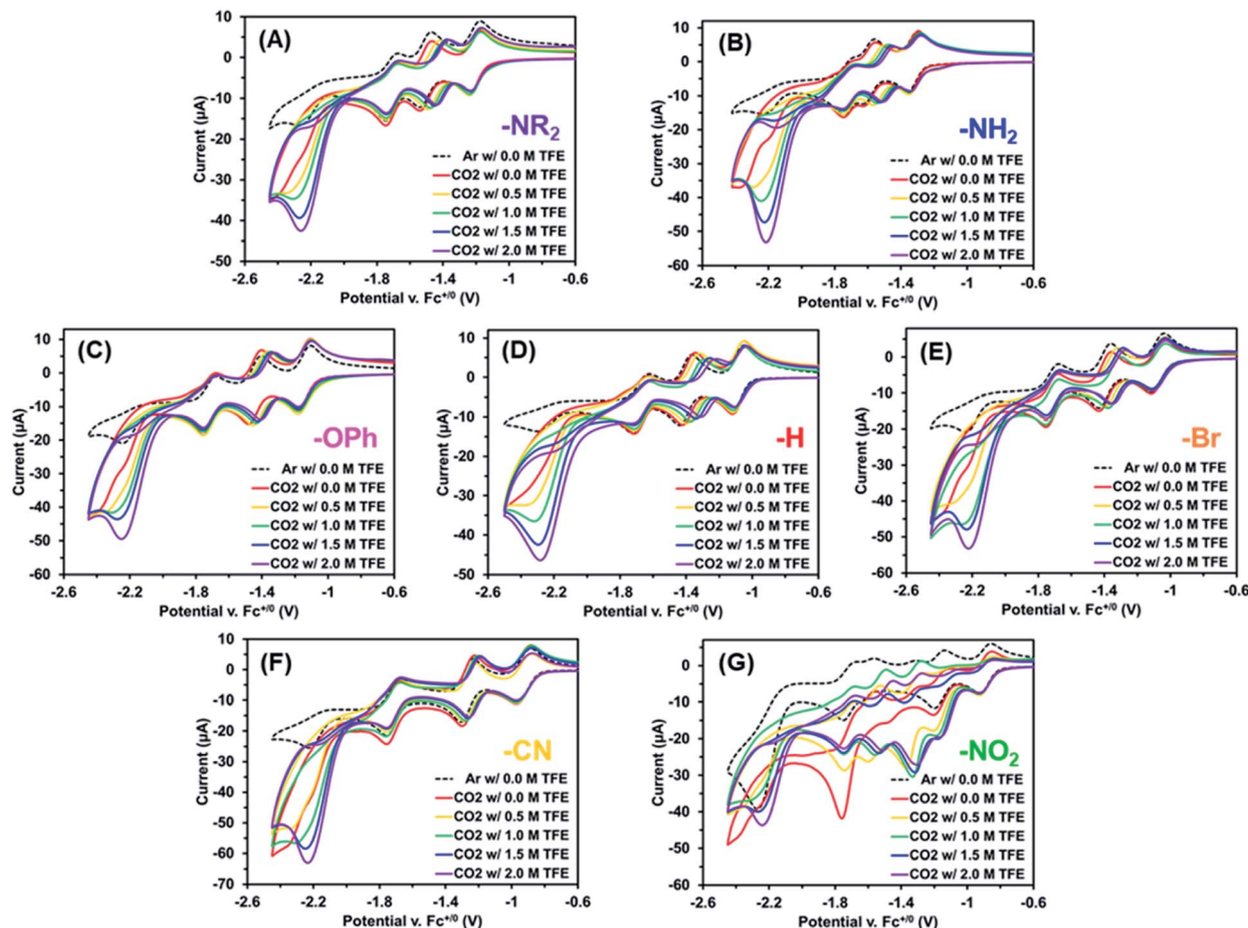


Fig. 3 CV current enhancement plots as a function of 2,2,2-trifluoroethanol (TFE) concentration for  $\text{Re}(\text{bpy-C2-NPDI-R})$  derivatives, where  $\text{R} = -\text{NR}_2$  (A),  $\text{R} = -\text{NH}_2$  (B),  $\text{R} = -\text{OPh}$  (C),  $\text{R} = -\text{H}$  (D),  $\text{R} = -\text{Br}$  (E),  $\text{R} = -\text{CN}$  (F), and  $\text{R} = -\text{NO}_2$  (G). TFE was incrementally added at 0 M (red), 0.5 M (yellow), 1.0 M (green), 1.5 M (blue), and 2 M (purple). All measurements were recorded at  $100 \text{ mV s}^{-1}$  under an atmosphere of  $\text{CO}_2$ , in DMF with 0.1 M  $\text{TBAPF}_6$  supporting electrolyte (WE = glassy carbon, CE = Pt-wire, RE =  $\text{Ag}/\text{AgCl}$ , and  $\text{Fc}^{+/0}$  as internal reference standard). Note, the observed current enhancement at  $E_p = -1.35 \text{ V}$  (vs.  $\text{Fc}^{+/0}$ ) for  $\text{R} = -\text{NO}_2$  (G) results from an *in situ* electrochemical conversion to  $\text{R} = -\text{NH}_2$ .

enhancement increase under the fourth reduction event (Fig. 3). We further note that incremental addition of TFE also caused the  $\text{NPDI}^{2-/1-}$  redox couple to gradually shift to more positive potentials. This behavior has been modeled for similar rylene diimide materials as a  $2\text{H}^+/2\text{e}^-$  proton-coupled electron-transfer process,<sup>65</sup> whereby the NPDI imide oxygens are protonated by the proton-source.<sup>66,67</sup> While the collected CV data was not obtained under steady-state conditions (*i.e.* plateau current),<sup>68</sup> modelling these measured CV current enhancements as  $\text{CO}_2$  reduction variables can qualitatively describe the effects of catalyst and proton-source as first-order and second-order rate-dependent variables, respectively. In other words, these data imply that one  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyad, with the assistance of two proton-source molecules, can enable electrocatalytic  $\text{CO}_2$  conversion.<sup>45</sup> The obvious exception to these generalized electrochemical trends is the  $-\text{NO}_2$  dyad (Fig. 3G). An in-depth evaluation of the electrochemical behaviour of  $-\text{NO}_2$  under argon and  $\text{CO}_2$  is provided in the ESI (see Section VIII, Fig. S84–S86<sup>†</sup>). Thorough analysis of this data strongly suggests that  $-\text{NO}_2$  is converted *in situ* to  $-\text{NH}_2$  under

reducing conditions; as such, it is difficult to establish meaningful trends for  $-\text{NO}_2$  (*vide infra*).

### $\text{CO}_2$ electro-/photocatalysis

The electrocatalytic  $\text{CO}_2$  reduction abilities of these  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads were evaluated using controlled potential electrolysis (CPE). All experiments were performed in DMF (with 2 M TFE) using our previously described two-compartment H-cell.<sup>45,69</sup> At an applied potential ( $E_{\text{app}}$ ) of  $-1.8 \text{ V}$  (Fig. S75<sup>†</sup>), all  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads (except  $-\text{CN}$ ) achieved comparable turnover numbers of CO ( $\text{TON}_{\text{CO}} = 21\text{--}25$ ) and faradaic efficiencies ( $\text{FE}_{\text{CO}} = 87\text{--}99\%$ ) after 6 hours of electrocatalysis. The  $-\text{CN}$  derivative, on the other hand, attained about half the performance ( $\text{TON}_{\text{CO}} = 13$ ) at a slightly lower  $\text{FE}_{\text{CO}}$  (87%). This drop in performance is consistent with decreased efficacy of the electron-reservoir effect due to the increased electron-affinity of NPDI-CN (Fig. S77<sup>†</sup>). Unlike the electronically modified  $\text{Re}(4,4'\text{-R-bpy})$  series reported by Kubiak *et al.*, where electrocatalytic  $\text{CO}_2$ -to-CO conversion efficiency was highly dependent on the nature of the R-group,<sup>11</sup> all  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads achieved



high  $FE_{CO}$  at an overpotential that is  $\sim 300$  mV lower than the measured  $E_{cat/2} \approx -2.1$  V (Fig. S67†). It should be noted, however, that when an  $E_{app1} = -1.7$  V was used for CPE (Fig. S76†), electrocatalytic  $CO_2$  reduction was essentially shut-off for all **Re(bpy-C2-NPDI-R)** dyads ( $TON_{CO} \leq 6$ ). This could indicate that altering the electronic properties of the electron-reservoir may not be the most effective strategy towards further lowering the overpotentials required to enable electrocatalytic  $CO_2$  reduction.

In our previous report, we noticed that the use of blue light during CPE experiments significantly increased the rates of CO production for  $-H$ .<sup>45</sup> To determine the origin of this  $TON_{CO}$  enhancement,<sup>28,70,71</sup> we next turned our attention towards the photocatalytic  $CO_2$  reduction capabilities of these **Re(bpy-C2-NPDI-R)** dyads. Following established literature protocols,<sup>31,34–37</sup> each dyad (30  $\mu M$ ) was dissolved in a (5 : 1) DMF: triethanolamine (TEOA) mixture containing sacrificial reducing agent 1,3-dimethyl-2-phenyl-2,3-dihydro-1H-benzo[d]imidazole (BIH; 3 mM). The glass vials were sealed with rubber septa, sparged with  $CO_2$ , and then irradiated with blue light ( $\lambda = 470 \pm 30$  nm; 4  $mW\ cm^{-2}$ ) for 24 hours (Table 1).

Over the 24 h testing period, the **Re(bpy-C2-NPDI-R)** dyads all showed good activity for  $\sim 9$  hours, after which CO production would level-off for the remainder of the experiment (Fig. S78†). The benchmark dyad,  $-H$ , achieved a  $TON_{CO}$  of  $57 \pm 1$  with a selectivity for CO of 97%. The  $-Br$  ( $TON_{CO} = 61 \pm 5$ ) and  $-OPh$  ( $TON_{CO} = 59 \pm 6$ ) dyads achieved the same performance with roughly the same CO selectivity ( $\geq 95\%$ ). The  $-CN$  derivative was completely inactive for  $CO_2$  photocatalysis under these conditions. Interestingly, despite being the most EWG, the  $-NO_2$  dyad achieved the second best  $TON_{CO}$  ( $134 \pm 15$ ) with a very high CO selectivity of 98%. When EDGs were functionalized on NPDI, the  $CO_2$ -to-CO production and selectivity was improved to ( $TON_{CO} = 86 \pm 8$ ) for  $-NR_2$  and ( $TON_{CO} = 234 \pm 13$ ) for  $-NH_2$ . We note that the photocatalytic performance of  $-NH_2$  was not greatly improved over the 24 h testing period by replenishing

both BIH and  $CO_2$  in 6 h intervals ( $TON_{CO} = 294$ ; Fig. S79†), suggesting that depletion of substrate was not the limiting factor for  $TON_{CO}$ .

To confirm the importance of each component in the photocatalytic  $CO_2$  reduction setup, various control experiments were conducted. As expected, the omission of **Re(bpy-C2-NPDI-R)** dyads or  $CO_2$  from the setup stopped the production of CO. When sacrificial reducing agent BIH was excluded (Table S2†), the  $TON_{CO}$  was decreased by at least two-fold for all dyads. This result implies that while TEOA alone can simultaneously act as the proton-source and the sacrificial electron-donor,<sup>20</sup> BIH is more efficient at reductively quenching the photoexcited dyads. When TEOA was replaced by TFE (Table S3†), the production of CO for all dyads was decreased almost four-fold, except for  $-NH_2$  ( $TON_{CO} = 143$ ). Overall, this result points towards the utility of TEOA as a sacrificial electron-donor, as well as the importance of forming **Re(bpy)**-adducts during  $CO_2$  reduction, as seen in previous literature examples.<sup>5,26–31</sup> When the irradiation source was switched to a green light LED array ( $\lambda = 525 \pm 32$  nm; 1.9  $mW\ cm^{-2}$ ), the measured  $TON_{CO}$  decreased at least five-fold for all **Re(bpy-C2-NPDI-R)** dyads (Table S4†). This significant drop in performance could be the result of either inefficient photoexcitation pathways in the dyad<sup>40–44</sup> and/or the elimination of a photo-assisted CO cleavage process.<sup>70,71</sup> Lastly, the importance of tethering the NPDI-moiety to the **Re(bpy)**-moiety was demonstrated by combining  $N_3$ -C2-NPDI-R with **Re(bpy)**, where it was observed that all samples obtained the same performance as **Re(bpy)** alone (Table S5†). We further showed that all  $N_3$ -C2-NPDI-R precursors were essentially inactive for  $CO_2$  conversion under the optimized photocatalysis conditions ( $TON_{CO} < 3$ ; Table S6†).

While these control experiments clearly highlight the necessity of each component in the photocatalytic  $CO_2$  reduction process, it does not fully account for the performance differences of each **Re(bpy-C2-NPDI-R)** dyad. Therefore, based on the photocatalytic  $CO_2$  reduction results, the **Re(bpy-C2-NPDI-R)** dyads may be grouped together in three categories: (i) standard catalysts (weak EWGs/EDGs =  $-H$ ,  $-Br$ , and  $-OPh$ ), (ii) inactive catalysts (strong EWGs =  $-CN$ ), and (iii) top-performing catalysts (strong EDGs =  $-NR_2$  and  $-NH_2$ ). Note, the  $-NO_2$  dyad can also be classified as a top-performing catalyst because  $-NO_2$  undergoes an *in situ* transition to  $-NH_2$  under photocatalytic  $CO_2$  reduction conditions (see ESI Section VIII for more details†).

Table 1 Optimized photocatalytic  $CO_2$  reduction of **Re(bpy-C2-NPDI-R)**

R =	$TON_{CO}^a$	$TON_{H_2}^a$	CO : H <sub>2</sub> <sup>b</sup>
$-H$	$57 \pm 1$	$1.8 \pm 1$	97 : 3
$-CN$	<1	<1	—
$-NO_2^c$	$137 \pm 15$	$3 \pm 0.6$	98 : 2
$-Br$	$61 \pm 5$	$2.5 \pm 1$	96 : 4
$-NR_2$	$86 \pm 8$	$1.4 \pm 1$	98 : 2
$-OPh$	$59 \pm 6$	$3.6 \pm 2$	95 : 5
$-NH_2$	$234 \pm 13$	$2.0 \pm 0.1$	>99 : 1

<sup>a</sup> Calculated based on bulk catalyst concentration from quadruplicate trials. <sup>b</sup> No other gaseous (*i.e.*  $CH_4$ ) or liquid (*i.e.*  $HCOO^-$ ) products were detected. <sup>c</sup> **Re(bpy-C2-NPDI-NO<sub>2</sub>)** is *in situ* converted to **Re(bpy-C2-NPDI-NH<sub>2</sub>)**.

## Mechanistic investigation

During electro-/photocatalytic  $CO_2$  reduction testing of these **Re(bpy-C2-NPDI-R)** dyads, a series of dramatic colour changes were observed. Prior to light irradiation, the absorption profile of all **Re(bpy-C2-NPDI-R)** dyads were unchanged by the addition of both TEOA and BIH, confirming that neither reagent reduces **Re(bpy-C2-NPDI-R)** immediately. After sparging with  $CO_2$ , the samples were then subjected to blue light and the progression of colour changes was monitored periodically by UV-vis-NIR spectroscopy (Fig. S80†). Each **Re(bpy-C2-NPDI-R)** dyad underwent a transition from their initial colour to either a green ( $-H$ ,



-Br, and -OPh), a dark blue (-CN), or a beige (-NO<sub>2</sub>, -NR<sub>2</sub>, and -NH<sub>2</sub>) colour that was found to persist throughout the remainder of catalysis. Photoluminescence spectroscopy revealed that the photoexcited state of all **Re(bpy-C2-NPDI-R)** dyads can be reductively quenched by both TEOA and BIH (Fig. S81†). Thus, to gain further insight into the photoelectrochemical processes that were occurring during CO<sub>2</sub> reduction catalysis, UV-vis-nIR and FTIR spectroelectrochemistry (SEC) experimentation was conducted. UV-vis-nIR and FTIR SEC data was collected by monitoring an air-free **Re(bpy-C2-NPDI-R)** dyad solution that was held at a constant  $E_{\text{appl}}$  (where Red1, Red2, and Red3 correspond to the NPDI<sup>•-0</sup>, NPDI<sup>2-/-</sup>, and bpy<sup>•-0</sup> reductions, respectively).

The UV-vis-nIR SEC data of each **Re(bpy-C2-NPDI-R)** dyad (Fig. 4) correlates very well with what was observed when we periodically monitored our photocatalytic experiments. When using an  $E_{\text{appl}} = \text{Red1}$ , the  $\lambda_{\text{max}}$  of all **Re(bpy-C2-NPDI-R)** dyads was bathochromically shifted, the relative molar absorptivity of this  $\lambda_{\text{max}}$  was stronger, and some new vibrational fine-structure was also observed between 800–1000 nm. While the nature of installed R-group influences the position of the  $\lambda_{\text{max}}$  and the absorption fine-structure, all these absorption features are

consistent with selective formation of NPDI<sup>•-0</sup>.<sup>43,44</sup> When the  $E_{\text{appl}}$  is switched to Red2, the spectral features of NPDI<sup>•-0</sup> are rapidly depleted and replaced by shifted broad-band absorption peak(s). These spectral features are commonly associated with NPDI<sup>2-/-</sup>,<sup>67</sup> where the vibronic structure of the NPDI<sup>2-/-</sup> absorption profile is once again influenced by the nature of installed R-group. With respect to -H (Fig. 4D), the incorporation of HOMO-modifying EDGs (*i.e.* -NR<sub>2</sub>, -NH<sub>2</sub>, and -OPh) resulted in minimal changes to the NPDI<sup>2-/-</sup> absorption profile shape (Fig. 4A–C). Conversely, when LUMO-modifying EWGs (*i.e.* -Br and -CN) are used, the vibronic structure of NPDI<sup>2-/-</sup> is significantly different (Fig. 4E and F). When the  $E_{\text{appl}}$  was changed to Red3, no other significant spectral changes were observed. This result is unsurprising given the differences in molar absorptivity of the NPDI and Re(bpy)-moieties.<sup>57,72</sup> Except for -NO<sub>2</sub> (Fig. 4G), all **Re(bpy-C2-NPDI-R)** dyads displayed very similar spectral transitions. In-depth analysis of the UV-vis-nIR SEC data for -NO<sub>2</sub> (Fig. S86†) shows that -NO<sub>2</sub> is *in situ* converted to -NH<sub>2</sub> under the conditions necessary for CO<sub>2</sub> reduction catalysis. These results help confirm why -NO<sub>2</sub> served as an efficient CO<sub>2</sub> reduction photocatalyst despite having a stronger EWG than the totally inactive -CN dyad derivative.

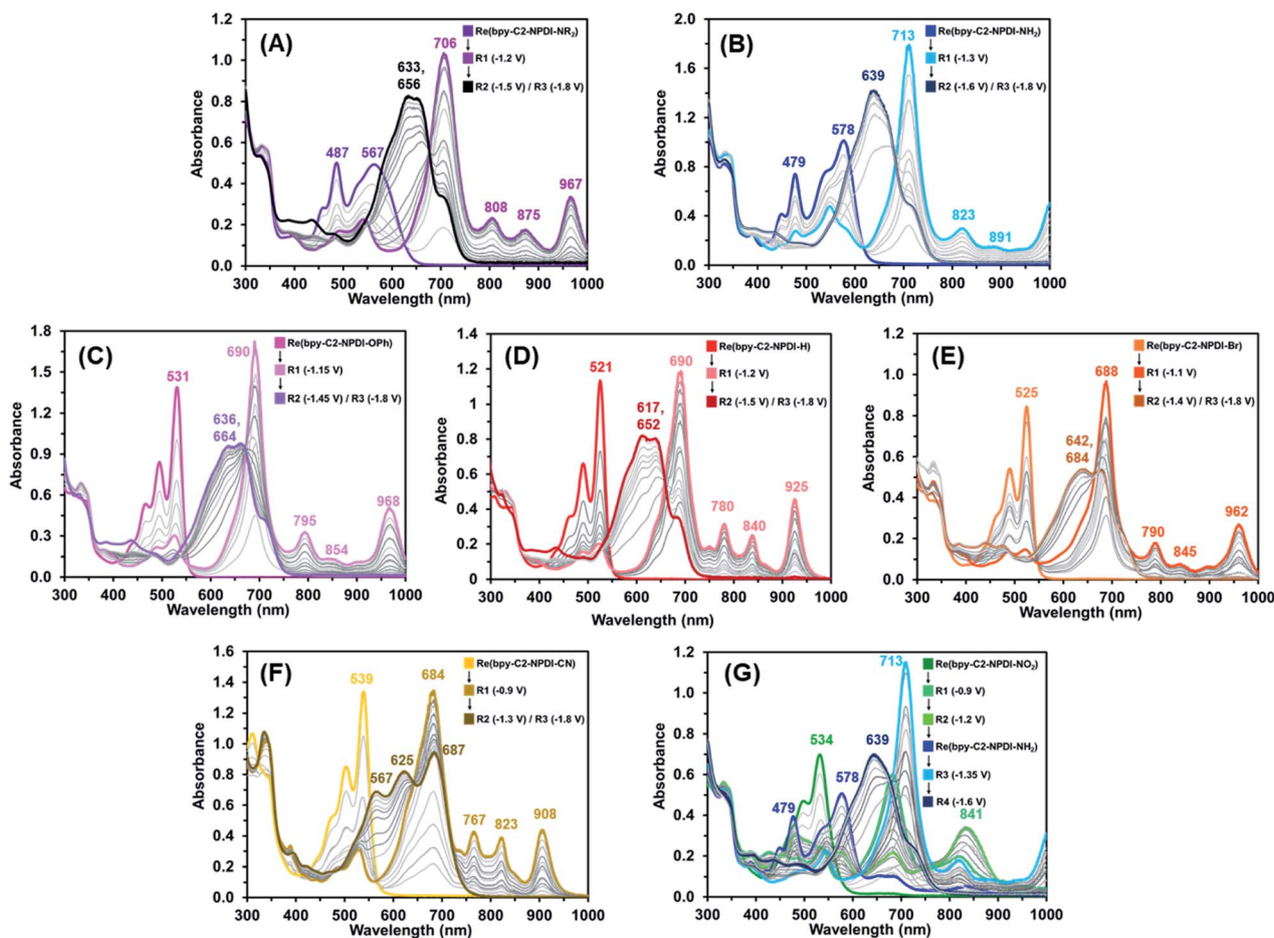


Fig. 4 UV-vis-nIR SEC absorbance spectra of **Re(bpy-C2-NPDI-R)** derivatives, where R = -NR<sub>2</sub> (A), R = -NH<sub>2</sub> (B), R = -OPh (C), R = -H (D), R = -Br (E), R = -CN (F), and R = -NO<sub>2</sub> (G). All experiments were performed in DMF with 0.1 M TBAPF<sub>6</sub> supporting electrolyte (WE = Pt-mesh, CE = Pt-wire, pseudo-RE = Ag-wire).



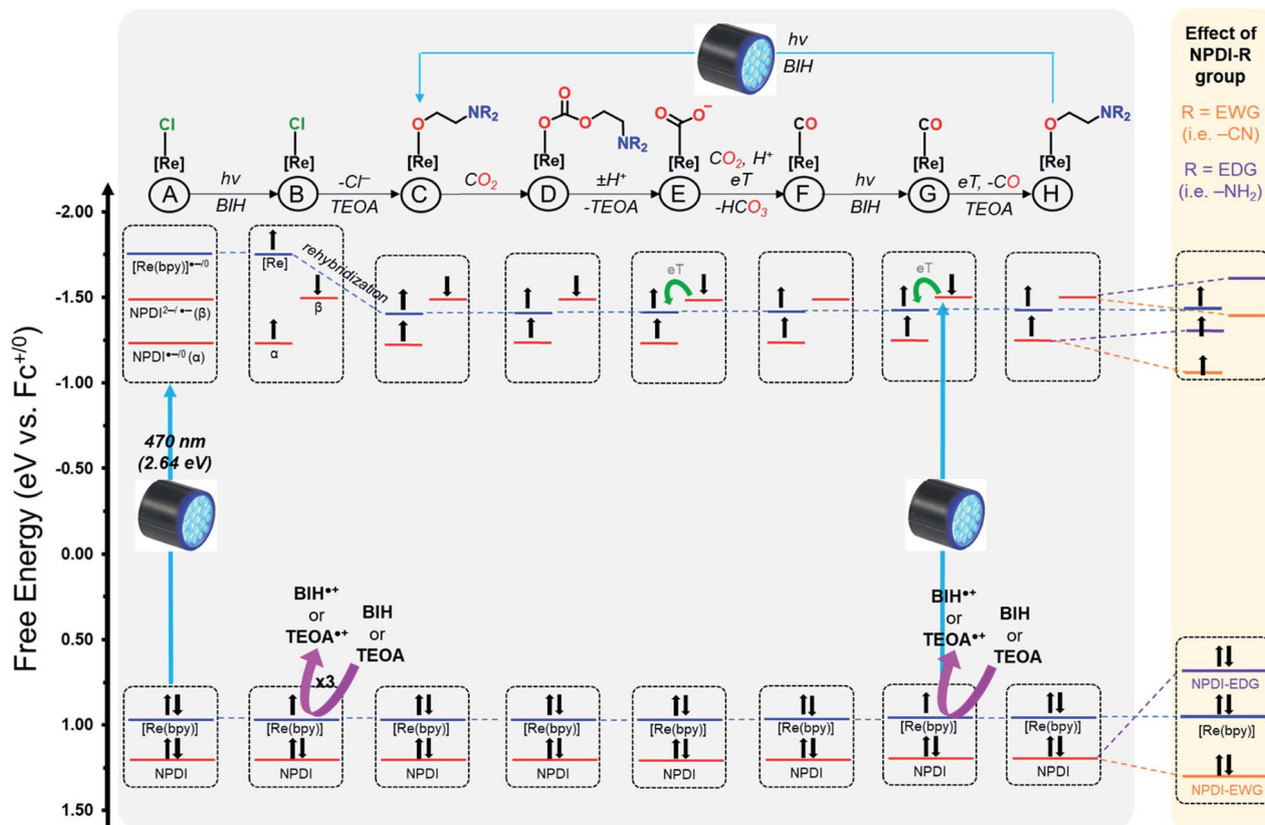


Fig. 5 Molecular orbital description for the proposed photocatalytic  $\text{CO}_2$  reduction mechanism of  $\text{Re}(\text{bpy-C2-NPDI-R})$ , where  $\text{Re}(\text{bpy})$  orbitals are shown in blue and NPDI-R orbitals are shown in red. The influence of electron withdrawing groups (EWGs, orange) and electron donating groups (EDGs, purple) on NPDI-R molecular orbitals is highlighted (right). Both BIH and TEOA are depicted as sacrificial electron donors that reductively quench the photoexcited electrons.

Moving onto the FTIR SEC data, the behavior of all  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads were nearly identical (Fig. S58–S63<sup>†</sup>). While no spectral changes were detected at  $E_{\text{appl}} = \text{Red1}$  and  $\text{Red2}$ , significant shifts in  $\nu_{\text{CO}}$  were observed at  $E_{\text{appl}} = \text{Red3}$ . At  $\text{Red3}$ , the  $\text{Re}(\text{bpy})$ -moiety of these dyads is formally reduced by one-electron [ $\text{Re}^{\text{I}}(\text{bpy}^{\cdot-}\text{-C2-NPDI}^{2-}\text{-R})$ ]. The added electron density at the  $\text{Re}(\text{bpy})$ - moiety results in a lowering of  $\nu_{\text{CO}}$  from 1895, 1915, and 2019  $\text{cm}^{-1}$  to roughly 1865, 1885, and 1995  $\text{cm}^{-1}$ , respectively. Over time, an equilibration process occurs whereby electron-density is shifted from  $\text{bpy}^{\cdot-}$  to  $\text{Re}$ , and leads to  $\text{Re-Cl}$  dissociation [ $\text{Re}^0(\text{bpy-C2-NPDI}^{2-}\text{-R})$ ].<sup>58</sup> This crucial process generates a 5-coordinate  $\text{Re}$  metal-center and can be characterized by a  $\Delta\nu_{\text{CO}}$  to 1843, 1862, and 1978  $\text{cm}^{-1}$ . Although this  $\text{Cl}$ -dissociation process was more readily observed for dyads bearing EWGs ( $-\text{CN}$ ,  $-\text{NO}_2$ , and  $-\text{Br}$ ) than it was for dyads bearing EDGs ( $-\text{NR}_2$ ,  $-\text{NH}_2$ , and  $-\text{OPh}$ ), it was still detected to some degree for all  $\text{Re}(\text{bpy-C2-NPDI-R})$  catalysts.

The similarities between our SEC data and the results reported for other  $\text{Re}(\text{bpy})$  and  $\text{Ru}^{\text{II}}\text{-Re}^{\text{I}}$  catalysts suggests that our  $\text{Re}(\text{bpy-C2-NPDI-R})$  systems likely operate *via* similar photocatalytic  $\text{CO}_2$  reduction mechanisms.<sup>5,25–31</sup> It should be noted, however, that the photocatalytic  $\text{CO}_2$  reduction mechanism for  $\text{Re}(\text{bpy})$  is still heavily debated.<sup>18–31,73</sup> As such, the goal of our proposed mechanism was not to precisely determine the exact

identity, electron-spin configuration (singlet *vs.* triplet), and/or the rate dynamics of  $\text{eT}$  for all photocatalytic  $\text{CO}_2$  reduction intermediates. Instead, we focus on developing a molecular orbital (MO) description of these  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads that helps account for the observed differences in photocatalytic  $\text{CO}_2$  reduction performance (Fig. 5). The presented mechanism was modeled after the benchmark dyad,  $-\text{H}$ .

By CV, it was elucidated for all  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads (except  $-\text{NH}_2$  and  $-\text{NR}_2$ ) that the HOMO is  $\text{Re}(\text{bpy})$ -based and the LUMO is NPDI-based (Fig. 5A). Consequently, the first two photoreductions most likely result from a  $\text{Re}-\pi^*$  intersystem crossing (ISC) process; however, a direct NPDI-based  $\pi-\pi^*$  transition may also be possible.<sup>43,44</sup> The third photoreduction likely occurs either *via* direct  $\text{Re}(\text{bpy})$  <sup>3</sup>MLCT<sup>19</sup> or  $\text{eT}$  from a photoexcited state of  $\text{NPDI}^{2-}$ .<sup>43,44</sup> Following the formal three electron reduction of the  $\text{Re}(\text{bpy-C2-NPDI-R})$  dyads (Fig. 5B), the next step is  $\text{Re-Cl}$  dissociation which generates a 5-coordinate  $\text{Re}$  metal-center whose axial position subsequently forms an adduct with TEOA.<sup>20,74</sup> Another consequence of the chloro-dissociation step is that the  $\text{Re}$  metal-center undergoes a rehybridization process that lowers the overall energy of the  $\text{Re}(\text{bpy})$ -based MOs.<sup>45,75</sup> Following the formation of the TEOA- $\text{Re}(\text{bpy})$  adduct, it is possible for  $\text{CO}_2$  insertion to occur without direct  $\text{eT}$  from the catalyst center (Fig. 5D).<sup>20,27–30</sup> Protonation of the



resulting carbonate intermediate induces a reorganization process that releases TEOA and forms a  $\text{Re-CO}_2^{\cdot-}$  species (Fig. 5E).<sup>22</sup> Due to the presence of  $^{13}\text{CO}$  and  $\text{H}^{13}\text{CO}_3^-$  in the  $^{13}\text{C}$   $\{^1\text{H}\}$  NMR spectrum after blue light irradiation (Fig. S83†), it is postulated that these **Re(bpy-C2-NPDI-R)** dyads operate *via* a BIH-mediated disproportionation reaction between  $\text{Re-CO}_2^{\cdot-}$  and another equivalent of  $\text{CO}_2$  that liberates  $\text{HCO}_3^-$ ,<sup>25</sup> rather than a proton-coupled electron transfer process from  $\text{NPDI}^{2-}$  to the **Re(bpy)**-moiety that liberates  $\text{OH}^-$ .<sup>18</sup> From there, reductive quenching of a photoexcited electron restores  $\text{NPDI}^{2-}$  (Fig. 5G) and the ensuing transfer of this electron to the **Re(bpy)**-moiety produces CO, as well as opens a coordination site for TEOA (Fig. 5H).<sup>21</sup> The photocatalytic cycle is completed by the photoexcitation and reductive quenching of the electron to regenerate  $\text{NPDI}^{2-}$ .

Based on photocatalytic  $\text{CO}_2$  reduction performance, the catalysts were loosely grouped into three categories: (i) standard catalysts ( $-\text{H}$ ,  $-\text{Br}$ , and  $-\text{OPh}$ ), (ii) inactive catalysts ( $-\text{CN}$ ), and (iii) top-performing catalysts ( $-\text{NR}_2$  and  $-\text{NH}_2$ ). Looking at the proposed photocatalytic  $\text{CO}_2$  reduction mechanism, it is also possible to map out the effects of EWGs and EDGs on the provided MO description of these **Re(bpy-C2-NPDI-R)** dyads (Fig. 5, highlighted in yellow). In the case of  $-\text{Br}$  and  $-\text{OPh}$ , the overall influence of these R-group does not appear to change the eT dynamics of the **Re(bpy-C2-NPDI-R)** dyad (with respect to  $-\text{H}$ ). The HOMO–LUMO transition ( $\text{Re}-\pi^*$ ) is identical and the relative shift(s) of the  $\text{NPDI}^{2-}$  energy level does not significantly alter the electronic driving-force of eT between  $\text{NPDI}^{2-}$  and **Re(bpy)**. In the case of  $-\text{CN}$ , the fixation of that EWG on NPDI served to lower the energy of the NPDI-based HOMO and LUMOs. While the net result of this transformation retains the original HOMO–LUMO transition ( $\text{Re}-\pi^*$ ), it appears to lower the energy level of  $\text{NPDI}^{2-}$  enough to effectively prevent eT from  $\text{NPDI}^{2-}$  to the **Re(bpy)**-moiety, thus shutting down catalysis. Conversely, the installation of strong EDGs ( $-\text{NR}_2$  and  $-\text{NH}_2$ ) causes the energy levels of the NPDI-based HOMO and LUMOs to increase. The overall result of this transformation not only changes the HOMO–LUMO transition to an exclusively NPDI-based process ( $\pi-\pi^*$ ), but it also increases the driving-force for eT from  $\text{NPDI}^{2-}$  to **Re(bpy)**. The sum of these two effects together lead to improved dyad eT dynamics, thereby enhancing photocatalytic  $\text{CO}_2$  reduction (with respect to  $-\text{H}$ ).

It should be noted that another feasible explanation for the improved performance of the amino-functionalized dyads, in particular  $-\text{NH}_2$ , is the possibility of second-sphere H-bonding effects.<sup>15,18</sup> Previously we calculated the optimized geometries of various intermediates during  $\text{CO}_2$  catalysis for the  $-\text{H}$  dyad.<sup>45</sup> It was shown in this study that, although the NPDI was initially folded over the **Re(bpy)**-moiety, reduction of the  $-\text{H}$  dyad caused the two moieties to extend away from one another (likely due to coulombic repulsion effects). By analogy, the amino-functionalized NPDI-R bay position would most likely also be extended away from the **Re(bpy)**-moiety during  $\text{CO}_2$  photocatalysis. While in this case the distance between the catalyst center and the amino-groups of  $-\text{NR}_2$  and  $-\text{NH}_2$  make it unlikely that second-sphere H-bonding effects are involved in  $\text{CO}_2$  photocatalysis, they can't be conclusively ruled out at this time.

If nothing else, the synthetic versatility of the NPDI chromophore means that future iterations of the **Re(bpy-C2-NPDI-R)** motif could incorporate proximal second-sphere H-bonding groups as a means to further improve  $\text{CO}_2$  conversion performance.

## Conclusions

In conclusion, we present the synthesis and full characterization of six new **Re(bpy-C2-NPDI-R)** supramolecular dyad materials (where  $\text{R} = -\text{Br}$ ,  $-\text{CN}$ ,  $-\text{NO}_2$ ,  $-\text{OPh}$ ,  $-\text{NH}_2$ , or  $-\text{NR}_2$ ). The installation of R-groups on NPDI altered the optoelectronic properties of these dyads, as well as impacted the photocatalytic  $\text{CO}_2$  reduction performance. Relative to the benchmark **Re(bpy-C2-NPDI-H)** dyad ( $\text{TON}_{\text{CO}} = 57$ ), the incorporation of EDGs (*i.e.*  $-\text{NH}_2$ ) led to an over four-fold improvement in photocatalytic  $\text{CO}_2$  reduction performance ( $\text{TON}_{\text{CO}} = 234$ ) while strong EWGs (*i.e.*  $-\text{CN}$ ) resulted in complete deactivation of the dyads. Despite being the most electron-withdrawing, the  $-\text{NO}_2$  functionalized NPDI was among the top performing  $\text{CO}_2$  reduction photocatalysts ( $\text{TON}_{\text{CO}} = 137$ ), making it an outlier to the proposed trend. Through CV and UV-vis-nIR SEC experimentation, it was elucidated that  $-\text{NO}_2$  undergoes an *in situ* conversion to  $-\text{NH}_2$ , thereby forming a different dyad that is responsible for catalysis. A photocatalytic  $\text{CO}_2$  reduction mechanism is proposed for these dyads, where EDGs served to accelerate  $\text{CO}_2$  reduction rates by simultaneously changing the HOMO–LUMO excitation pathway and by increasing the electronic driving-force of intramolecular electron transfer from  $\text{NPDI}^{2-}$  to **Re(bpy)**. Conversely, EWGs shifted the LUMO energy levels of NPDI to the point where photocatalysis is shut down because there is no electronic driving force for eT between  $\text{NPDI}^{2-}$  and **Re(bpy)**. This study clearly highlights the importance of evaluating structure–property relationships to develop and optimize the future design of new supramolecular dyad photocatalysts.

## Author contributions

JDBK performed all experimental work and data analysis and prepared the manuscript. GCW directed the project and provided resources. WEP co-directed the project and provided resources.

## Conflicts of interest

There are no conflicts to declare.

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## Notes and references

- 1 D. W. Keith, G. Holmes, D. S. Angelo and K. Heidel, *Joule*, 2018, **2**, 1573–1594.
- 2 P. De Luna, C. Hahn, D. Higgins, S. A. Jaffer, T. F. Jaramillo and E. H. Sargent, *Science*, 2019, **364**, eaav3506.
- 3 C. D. Windle and R. N. Perutz, *Coord. Chem. Rev.*, 2012, **256**, 2562–2570.
- 4 C. Costentin, M. Robert and J.-M. Savéant, *Chem. Soc. Rev.*, 2013, **42**, 2423–2436.
- 5 G. Sahara and O. Ishitani, *Inorg. Chem.*, 2015, **54**, 5096–5104.
- 6 H. Takeda, C. Cometto, O. Ishitani and M. Robert, *ACS Catal.*, 2017, **7**, 70–88.
- 7 R. Francke, B. Schille and M. Roemelt, *Chem. Rev.*, 2018, **118**, 4631–4701.
- 8 K. E. Dalle, J. Warnan, J. J. Leung, B. Reuillard, I. S. Karmel and E. Reisner, *Chem. Rev.*, 2019, **119**, 2752–2875.
- 9 F. Franco, C. Rettenmaier, H. S. Jeon and B. R. Cuenya, *Chem. Soc. Rev.*, 2020, **49**, 6884–6946.
- 10 J. Hawecker, J.-M. Lehn and R. Ziessel, *J. Chem. Soc., Chem. Commun.*, 1983, 536–538.
- 11 M. L. Clark, P. L. Cheung, M. Lessio, E. A. Carter and C. P. Kubiak, *ACS Catal.*, 2018, **8**, 2021–2029.
- 12 S. Sung, D. Kumar, M. Gil-Sepulcre and M. Nippe, *J. Am. Chem. Soc.*, 2017, **139**, 13993–13996.
- 13 E. Haviv, D. Azaiza-Dabbah, R. Carmieli, L. Avram, J. M. L. Martin and R. Neumann, *J. Am. Chem. Soc.*, 2018, **140**, 12451–12456.
- 14 A. N. Hellman, R. Haiges and S. C. Marinescu, *Dalton Trans.*, 2019, **48**, 14251–14255.
- 15 K. Talukdar, S. Sinha Roy, E. Amatya, E. A. Sleeper, P. Le Magueres and J. W. Jurss, *Inorg. Chem.*, 2020, **59**, 6087–6099.
- 16 M. R. Madsen, J. B. Jakobsen, M. H. Rønne, H. Liang, H. C. D. Hammershøj, P. Nørby, S. U. Pedersen, T. Skrydstrup and K. Daasbjerg, *Organometallics*, 2020, **39**, 1480–1490.
- 17 S. Lense, K. A. Grice, K. Gillette, L. M. Wolf, G. Robertson, D. McKeon, C. Saucedo, P. J. Carroll and M. Gau, *Organometallics*, 2020, **39**, 2425–2437.
- 18 J. Mukherjee and I. Siewert, *Eur. J. Inorg. Chem.*, 2020, **2020**, 4319–4333.
- 19 H. Takeda, K. Koike, H. Inoue and O. Ishitani, *J. Am. Chem. Soc.*, 2008, **130**, 2023–2031.
- 20 T. Morimoto, T. Nakajima, S. Sawa, R. Nakanishi, D. Imori and O. Ishitani, *J. Am. Chem. Soc.*, 2013, **135**, 16825–16828.
- 21 Y. Kou, Y. Nabetani, D. Masui, T. Shimada, S. Takagi, H. Tachibana and H. Inoue, *J. Am. Chem. Soc.*, 2014, **136**, 6021–6030.
- 22 T. W. Schneider, M. Z. Ertem, J. T. Muckerman and A. M. Angeles-Boza, *ACS Catal.*, 2016, **6**, 5473–5481.
- 23 P. Lang, R. Giereth, S. Tschierlei and M. Schwalbe, *Chem. Commun.*, 2019, **55**, 600–603.
- 24 J. H. Jo, S. Choi, H.-Y. Cheong, J. Y. Shin, C. H. Kim, D. W. Cho, H.-J. Son, C. Pac and S. O. Kang, *Chem.–Eur. J.*, 2020, **26**, 16733–16754.
- 25 Y. Tamaki, K. Koike, T. Morimoto and O. Ishitani, *J. Catal.*, 2013, **304**, 22–28.
- 26 Y. Kuramochi, O. Ishitani and H. Ishida, *Coord. Chem. Rev.*, 2018, **373**, 333–356.
- 27 K. Koike, D. C. Grills, Y. Tamaki, E. Fujita, K. Okubo, Y. Yamazaki, M. Saigo, T. Mukuta, K. Onda and O. Ishitani, *Chem. Sci.*, 2018, **9**, 2961–2974.
- 28 R. Kamata, H. Kumagai, Y. Yamazaki, G. Sahara and O. Ishitani, *ACS Appl. Mater. Interfaces*, 2019, **11**, 5632–5641.
- 29 Y. Yamazaki, K. Ohkubo, D. Saito, T. Yatsu, Y. Tamaki, S. Tanaka, K. Koike, K. Onda and O. Ishitani, *Inorg. Chem.*, 2019, **58**, 11480–11492.
- 30 K. Kamogawa, Y. Shimoda, K. Miyata, K. Onda, Y. Yamazaki, Y. Tamaki and O. Ishitani, *Chem. Sci.*, 2021, **12**, 9682–9693.
- 31 A. M. Cancelliere, F. Puntoriero, S. Serroni, S. Campagna, Y. Tamaki, D. Saito and O. Ishitani, *Chem. Sci.*, 2020, **11**, 1556–1563.
- 32 A. Gabrielsson, F. Hartl, H. Zhang, J. R. Lindsay Smith, M. Towrie, V. Antonin and R. N. Perutz, *J. Am. Chem. Soc.*, 2006, **128**, 4253–4266.
- 33 K. Kiyosawa, N. Shiraishi, T. Shimada, D. Masui, H. Tachibana, S. Takagi, O. Ishitani, D. A. Tryk and H. Inoue, *J. Phys. Chem. C*, 2009, **113**, 11667–11673.
- 34 C. D. Windle, M. W. George, R. N. Perutz, P. A. Summers, X. Z. Sun and A. C. Whitwood, *Chem. Sci.*, 2015, **6**, 6847–6864.
- 35 P. Lang, M. Pfrunder, G. Quach, B. Braun-Cula, E. G. Moore and M. Schwalbe, *Chem.–Eur. J.*, 2019, **25**, 4509–4519.
- 36 Y. Kuramochi, Y. Fujisawa and A. Satake, *J. Am. Chem. Soc.*, 2020, **142**, 705–709.
- 37 Y. Kuramochi and A. Satake, *Chem.–Eur. J.*, 2020, **26**, 16365–16373.
- 38 F. Franco, C. Cometto, C. Garino, C. Minero, F. Sordello, C. Nervi and R. Gobetto, *Eur. J. Inorg. Chem.*, 2015, **2015**, 296–304.
- 39 D. R. Case, A. Spear, A. F. Henwood, M. Nanao, S. Dampf, T. M. Korter, T. Gunnlaugsson, J. Zubieta and R. P. Doyle, *Dalton Trans.*, 2021, **50**, 3479–3486.
- 40 J. F. Martinez, N. T. La Porte and M. R. Wasielewski, *J. Phys. Chem. C*, 2018, **122**, 2608–2617.
- 41 J. F. Martinez, N. T. La Porte, S. Chaudhuri, A. Sinopoli, Y. J. Bae, M. Sohail, V. S. Batista and M. R. Wasielewski, *J. Phys. Chem. C*, 2019, **123**, 10178–10190.
- 42 J. F. Martinez, N. T. La Porte and M. R. Wasielewski, *J. Photochem. Photobiol., A*, 2019, **372**, 21–28.
- 43 S. Hedström, S. Chaudhuri, N. T. La Porte, B. Rudsteyn, J. F. Martinez, M. R. Wasielewski and V. S. Batista, *J. Am. Chem. Soc.*, 2017, **139**, 16466–16469.
- 44 N. T. L. Porte, J. F. Martinez, S. Hedström, B. Rudsteyn, B. T. Phelan, C. M. Mauck, R. M. Young, V. S. Batista and M. R. Wasielewski, *Chem. Sci.*, 2017, **8**, 3821–3831.
- 45 J. D. B. Koenig, Z. S. Dubrawski, K. R. Rao, J. Willkomm, B. S. Gelfand, C. Risko, W. E. Piers and G. C. Welch, *J. Am. Chem. Soc.*, 2021, **143**, 16849–16864.
- 46 C. Hansch, A. Leo and R. W. Taft, *Chem. Rev.*, 1991, **91**, 165–195.



- 47 J. R. Cann, C. Cabanetos and G. C. Welch, *Eur. J. Org. Chem.*, 2018, **2018**, 6933–6943.
- 48 A. D. Hendsbee, J.-P. Sun, W. K. Law, H. Yan, I. G. Hill, D. M. Spasyuk and G. C. Welch, *Chem. Mater.*, 2016, **28**, 7098–7109.
- 49 C. R. Harding, J. Cann, A. Laventure, M. Sadeghianlemraski, M. Abd-Ellah, K. R. Rao, B. S. Gelfand, H. Aziz, L. Kaake, C. Risko and G. C. Welch, *Mater. Horiz.*, 2020, **7**, 2959–2969.
- 50 M. Vespa, J. R. Cann, S. V. Dayneko, O. A. Melville, A. D. Hendsbee, Y. Zou, B. H. Lessard and G. C. Welch, *Eur. J. Org. Chem.*, 2018, **2018**, 4592–4599.
- 51 A. Call, M. Cibian, K. Yamamoto, T. Nakazono, K. Yamauchi and K. Sakai, *ACS Catal.*, 2019, 4867–4874.
- 52 X. Zhang, K. Yamauchi and K. Sakai, *ACS Catal.*, 2021, 10436–10449.
- 53 C. D. Sahm, G. M. Ucoski, S. Roy and E. Reisner, *ACS Catal.*, 2021, 11266–11277.
- 54 L. Hao, W. Jiang and Z. Wang, *Tetrahedron*, 2012, **68**, 9234–9239.
- 55 G. Li, Y. Zhao, J. Li, J. Cao, J. Zhu, X. W. Sun and Q. Zhang, *J. Org. Chem.*, 2015, **80**, 196–203.
- 56 D. H. Harris, S. Brixi, B. S. Gelfand, B. H. Lessard and G. C. Welch, *J. Mater. Chem. C*, 2020, **8**, 9811–9815.
- 57 S. Sato, Y. Matubara, K. Koike, M. Falkenström, T. Katayama, Y. Ishibashi, H. Miyasaka, S. Taniguchi, H. Chosrowjan, N. Mataga, N. Fukazawa, S. Koshihara, K. Onda and O. Ishitani, *Chemistry*, 2012, **18**, 15722–15734.
- 58 C. W. Machan, M. D. Sampson, S. A. Chabolla, T. Dang and C. P. Kubiak, *Organometallics*, 2014, **33**, 4550–4559.
- 59 N. Elgrishi, K. J. Rountree, B. D. McCarthy, E. S. Rountree, T. T. Eisenhart and J. L. Dempsey, *J. Chem. Educ.*, 2018, **95**, 197–206.
- 60 A. J. Bard and L. R. Faulkner, *Electrochemical Methods, Electrochemical Methods: Fundamentals and Applications*, Wiley, 2nd edn, 2008.
- 61 J.-M. Savéant, *Chem. Rev.*, 2008, **108**, 2348–2378.
- 62 C. Costentin and J.-M. Savéant, *ChemElectroChem*, 2014, **1**, 1226–1236.
- 63 S. Dey, M. E. Ahmed and A. Dey, *Inorg. Chem.*, 2018, **57**, 5939–5947.
- 64 M. E. Ahmed, A. Rana, R. Saha, S. Dey and A. Dey, *Inorg. Chem.*, 2020, **59**, 5292–5302.
- 65 C. Wiberg, M. Busch, L. Evenäs and E. Ahlberg, *Electrochim. Acta*, 2021, **367**, 137480.
- 66 E. Shirman, A. Ustinov, N. Ben-Shitrit, H. Weissman, M. A. Iron, R. Cohen and B. Rybtchinski, *J. Phys. Chem. B*, 2008, **112**, 8855–8858.
- 67 S. Seifert, D. Schmidt and F. Würthner, *Chem. Sci.*, 2015, **6**, 1663–1667.
- 68 E. S. Rountree, B. D. McCarthy, T. T. Eisenhart and J. L. Dempsey, *Inorg. Chem.*, 2014, **53**, 9983–10002.
- 69 J. D. B. Koenig, J. Willkomm, R. Roesler, W. E. Piers and G. C. Welch, *ACS Appl. Energy Mater.*, 2019, **2**, 4022–4026.
- 70 C. Cometto, L. Chen, P.-K. Lo, Z. Guo, K.-C. Lau, E. Anxolabéhère-Mallart, C. Fave, T.-C. Lau and M. Robert, *ACS Catal.*, 2018, 3411–3417.
- 71 S. Fernández, F. Franco, C. Casadevall, V. Martín-Diaconescu, J. M. Luis and J. Lloret-Fillol, *J. Am. Chem. Soc.*, 2020, **142**, 120–133.
- 72 S. S. Saund, M. A. Siegler and V. S. Thoi, *Inorg. Chem.*, 2021, **60**, 13011–13020.
- 73 A. C. Tsipis and A. A. Sarantou, *Dalton Trans.*, 2021, **50**, 14797–14809.
- 74 H. Koizumi, H. Chiba, A. Sugihara, M. Iwamura, K. Nozaki and O. Ishitani, *Chem. Sci.*, 2019, **10**, 3080–3088.
- 75 R. N. Schaugaard, K. Raghavachari and L. Li, *Inorg. Chem.*, 2018, **57**, 10548–10556.

