Electronic Effects of Substituents on fac-M(bpy-R)(CO)₃ (M = Mn, Re) Complexes for Homogeneous CO₂ Electroreduction

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Synthesis and characterization of 14 new 2,2′-bipyridine metal complexes fac-M(bpy-R)(CO)₃X (where M = Mn, X = Br or M = Re, X = Cl and R = -CF₃, -CN, -Ph, -PhOH, -NMe₂) are reported. The complexes have been characterized by NMR, IR spectroscopy and elemental analysis. Single crystal X-Ray diffraction structures have been solved for Re(dpbpy)(CO)₃Cl (dpbpy = 4,6-diphenyl-2,2′-bipyridine) and Re(hpbpy)(CO)₃Cl (hpbpy = 4-(2-hydroxy-phenyl)-6-phenyl-2,2′-bipyridine). Electrochemical behaviors of the complexes in acetonitrile under Ar and their catalytic performances for CO₂ reduction with added water and MeOH have been investigated by cyclic voltammetry and controlled potential electrolysis. The role of the substituents on the electrochemical properties and the related over potentials required for CO₂ transformation have been analyzed. The complexes carrying only electron withdrawing groups like -CF₃, -CN totally lose their catalytic activities toward CO₂ reduction, whereas the symmetric -NMe₂ substituted and push-pull systems (containing both -NMe₂ and -CF₃) still display electrocatalytic current enhancement under CO₂ atmosphere. The complexes carrying a phenyl or a phenol group in position 4 show catalytic behaviors similar to those of simple M-bpy systems. The only detected reduction product by GC analysis is CO: for example, fac-Re(bpy-4,4′-NMe₂)(CO)₃Cl gives CO with high faradic efficiency and a TON of 18 and 31, in absence of external proton source and with 5% MeOH, respectively. DFT calculations were carried out to highlight the electronic properties of the complexes; results are in agreement with experimental electrochemical data.

Keywords: CO₂ electroreduction, Mn complexes, Re complexes, DFT calculations, bipy ligands, homogeneous catalysis, electron-withdrawing, electron-donating
INTRODUCTION

Nowadays the CO₂ concentration in the atmosphere is continuously increasing alongside with the overall world energy demand. Converting carbon dioxide via electrochemical reduction into useful chemicals and fuels for energy storage is an attractive and promising approach. CO₂ reduction is a competition between its thermodynamic and kinetic: the one electron reduction in water occurs at a very negative potential (−1.90 V vs. SCE at pH 7) (Hammouche et al., 1988; Saveant, 2008) because it requires a drastic change in the geometry, from the linear CO₂ molecule to the bent CO₂⁻ radical anion. The reason of such high negative overpotential is due to the slow kinetics of the electron transfer, which is associated to the different geometries of the neutral and reduced species, respectively. However, the reduction reactions involving multiple electron transfers coupled with proton transfers provide a significant lowering of the thermodynamic barrier. To avoid the CO₂⁻ as intermediate and to lower the energy cost of the reduction process, key catalytic strategies have been developed with the aim of obtaining the various products selectively. The best electrocatalysts currently studied work at a potential 100 mV negative with respect to E° CO₂/P (where P generically indicates the reduction products, CO, HCOOH, HCHO, CH₃OH, CH₂) (Francke et al., 2018; Francke et al., 2018). Despite the numerous advantages of heterogeneous electrocatalysis (Sun et al., 2016; Rotundo et al., 2019a), clever integration with the homogeneous counterpart allows a rational design of the catalysts, by tuning both the metal center and/or the ligand. One of the greatest challenge of the homogeneous approach lies in the search of stability, durability and improved turnover number (TON) efficiencies (Grice, 2017; Takeda et al., 2017; Wang et al., 2019). Bipyridine transition metal complexes represent one of the most studied classes of molecular electrocatalysts since the 1980s (Stanbury et al., 2018). Bipyridine has been extensively studied as ligand in the field of electro and photo-catalysis because of the capability to store electrons and subsequently delocalize electronic density on its π orbitals (Vlček, 2002; Elgrishi et al., 2017). Both fac-Re(bpy)(CO)₂Cl and fac-Mn(bpy)(CO)₂Br are capable of reducing CO₂ to CO with high faradaic efficiency (Haweker et al., 1984; Bourrez et al., 2011). Comparing these two complexes, Mn exhibits a catalytic peak that is shifted more anodically (around 300 mV, Eₚ = −1.51 V vs. Ag/AgCl in MeCN) in comparison with the second reduction of Re analogous (Eₚ = −1.8 V vs. Ag/AgCl in MeCN). Another important difference is that usually the Mn-bpy complexes show their catalytic activities only in the presence of external proton sources. To better shed light on this uniqueness, our research group synthesized Mn(pdpbpy)(CO)₂Br (Franco et al., 2014, 2017), in which two pendant phenolic groups act as local proton source (Costentin et al., 2012), capable of reducing CO₂ even in anhydrous acetonitrile. In this case a considerable amount of HCOOH was also detected. Conversely the analogous complex in which the OH groups were replaced by methoxy groups did not show any catalytic activity without the addition of Brønsted acid.

The electrochemical behavior of the two well-known Re and Mn-bpy complexes could be reasonably altered by varying the bipyridine moiety, i.e., introducing electron-withdrawing and electron-donating groups (Machan et al., 2014; Walsh et al., 2015; Stanbury et al., 2018). Kubiak and his group investigated the effect of 4,4′-di-tert-butyl-2,2′-bipyridine (tBu₂-bpy) firstly on rhenium carbonyl complexes, secondarily on manganese (Smieja and Kubiak, 2010; Smieja et al., 2013). In other works they studied the role of the modification in the 6,6′ position of the bipyridine (Sampson et al., 2014; Sampson and Kubiak, 2016). A similar approach has already been applied to Mo and W-bpy complexes (Franco et al., 2017; Rotundo et al., 2018), to both Mn and Re-bpy complexes by some of us (Franco et al., 2017; Rotundo et al., 2019b) and in the current work.

The target of the modification is the reduction potential of the catalyst, namely E°_{cat}: usually a less negative E°_{cat} corresponds to a decreased rate of CO₂ conversion (Francke et al., 2018). Electronic properties of organic groups are commonly described by the inductive (+I) and mesomeric (+M) effects. Substituents with electron withdrawing groups like -CF₃ (strong –I effect) and -CN (weaker –I effect) and with electron donating groups, like -N(Me)₂ (strong +M effect), -Ph and -PhOH (weaker +M effect), were placed in the 4,4′, 4,6, and 5,5′ positions of the bipyridine ligand coordinated to the metals. Combining both push and pull effects in the so called “push-pull” system, an electronic gradient is forced through the bipyridine. In this paper we explore the electrocatalytic properties of novel Mn and Re bipy-type complexes, bearing 7 differently substituted ligands (Scheme 1). More generally electron-donating groups are expected to convey greater nucleophilicity to the metal-center, although catalysis should require higher overpotentials. DFT calculations have been used as complementary tool to better correlate experimental electrochemical data, whereas Controlled Potential Electrolysis (CPE) experiments are useful to elucidate the stability and durability of the catalysts in acetonitrile solutions.

RESULTS

Synthesis and Structure

Fourteen new 2,2′-bipyridine metal complexes fac-M(bpy-R)(CO)₂X where M = Mn, X = Br or M = Re, X = Cl are reported, namely Mn(bpy-4,4′-CF₃)(CO)₂Br (1a), Mn(bpy-5,5′-CF₃)(CO)₂Br (1b), Mn(bpy-4,4′-CN)(CO)₂Br (1c), Mn(bpy-4,4′-CF₃-NMe₂)(CO)₂Br (1d), Mn(dpbpy)(CO)₂Br (1e), Mn(hpbpy)(CO)₂Br (1f), Mn(bpy-4,4′-NMe₂)(CO)₂Br (1g), Re(bpy-4,4′-CF₃)(CO)₂Cl (2a), Re(bpy-5,5′-CF₃)(CO)₂Cl (2b), Re(bpy-4,4′-CN)(CO)₂Cl (2c), Re(bpy-4,4′-CF₃-NMe₂)(CO)₂Cl (2d), Re(dpbpy)(CO)₂Cl (2e), Re(hpbpy)(CO)₂Cl (2f), Re(bpy-4,4′-NMe₂)(CO)₂Cl (2g).

The ligands and the corresponding Mn and Re complexes have been synthesized according to the procedure reported in the experimental section. The complexes have been characterized by NMR, IR spectroscopy and elemental analysis. Single crystal X-Ray diffraction structures have been solved for 2e and 2f (for XRD data see Tables S1–S6).

The complex Re(dpbpy)(CO)₂Cl (2e) crystallized from both acetonitrile and benzene solutions by slow evaporation, forming prismatic orange platelets of a phase of the pure molecular product (structure A, Figure S1a) and a solvate with two benzene
molecules (structure B, Figure S1b). The first presents the monoclinic centrosymmetric P2_1/a space group (Figure S1c) while the second the monoclinic centrosymmetric P2/n space group (Figure S1d). The complex Re(hbpp)(CO)Br (2f) crystallized in dark by slow evaporation of solutions of toluene and benzene as yellow platelets and from ethyl acetate as orange prisms, both stable to air (Figure S2a). The crystal structure has been obtained from a platelet obtained from toluene solution and has monoclinic P2_1/n space group type (Figure S2b). The structures of both 2e and 2f present a quite distorted octahedral geometry around the rhenium center, as can be seen by the values in Table 1. The coordination bond of N1 is 0.1 Å longer than that of N2 (see Figure 1 for numeration) and a similar asymmetry can be observed in the coordination of CO in trans position to the nitrogens. This asymmetric coordination, also present in other terpyridine derivatives (Anderson et al., 1990; Civitello et al., 1993; Wang et al., 2013; Klemens et al., 2016) is completely different from the very symmetrically bonded bpy derivatives (with N-Re average distances equal in the two coordinating pyridyl rings and long 2.17 Å). At the same time, while most of the 2,2′-bipyridine derivatives of fac-Re(CO)3Cl complex unit are almost planar respect to the basal OC–Re–CO plane (Kurz et al., 2006; Smieja and Kubia, 2010; Bullock et al., 2012; Machan et al., 2014; Manbeck et al., 2015), in the case of 2e and 2f, the ligand is distorted outside this plane (see Table 1). This behavior can be detected in all the terpyridines, in which the third pyridine ring is not coordinated to the metal center and is equivalent to a phenyl ring, and in 6-phenyl substituted 2,2′-bipyridine derivatives. The reason of such distortion is clarified by considering the steric hindrance of the vicinal phenyl ring (the distance between the nearest CO group and the centroid of the phenyl ring in ortho to N1 is about 3.2 Å) that pushes up all the framework of the organic ligand, modifying the Re environment. This phenyl group is rotated to follow the shape of carbonyls with a torsion angle respect to the central pyridine ring of about 130°.

The distortion effects observed in the solid state are predicted also by molecular DFT calculations. Indeed, the optimized geometries of 2e and 2f perfectly overlap with the experimentally determined structures (Figure 1D and Figure S3), confirming that these anomalies originate from the coordination of 6-phenyl substituted 2,2′-bipyridine ligands to carbonyl complexes and are not induced by crystal packing contributions.

| Table 1 | Selected X-ray bond lengths [Å] and angles [°] for 2e and 2f. |
|---------|---------------------------------|
|          | 2e (structure A) | 2e (structure B) | 2f |
| Re–C1    | 1.906 (5)        | 1.919 (5)        | 1.919 (5) |
| Re–C2    | 1.936 (5)        | 1.952 (5)        | 1.935 (5) |
| Re–C3    | 1.890 (5)        | 1.933 (5)        | 1.898 (5) |
| Re–N1    | 2.160 (5)        | 2.177 (5)        | 2.164 (5) |
| Re–N2    | 2.220 (5)        | 2.233 (5)        | 2.214 (5) |
| N2–C13–C14–C15 | 125.66 (4) | 125.47 (4) | |
| C2O2–ph | 3.110 (5) | 3.239 (5) | 3.233 (5) |
| Re1–N2–C11 | 165.28 (6) | 162.38 (6) | 159.09 (6) |

The crystal packing of 2e (both structure A and structure B) and 2f is dominated by weak C–H···Cl, C–H···O and π···π stacking interactions. In the case of 2f, the presence of -OH group on the organic ligand interacting with the chloride induces the formation of hydrogen-bonded molecular chains (see Figure 1C and Figure S2c). For a more detailed analysis on the crystal packing consults the Supplementary Material (see Figures S4–S9).

Cyclic Voltammetry Under Ar

Cyclic Voltammetries (CVs) of all manganese complexes are reported in Figure 2. The CV of the Mn(bpy)(CO)3Br under our experimental conditions (in black) is included for comparison. This complex undergoes two successive irreversible reduction reactions and two reoxidation peaks (Bourrez et al., 2011). The first (E_p1 = −1.29 V vs. Ag/AgCl) and the second (E_p2 = −1.51 V vs. Ag/AgCl) reduction processes lead to the formation of the dimer and the mononuclear pentacoordinated anion species, respectively. Reoxidations of the pentacoordinated anion and of the dimer are located at −1.09 and −0.21 V vs. Ag/AgCl, respectively. The new synthesized complexes are supposed to display similar electrochemical behavior. Table 2 reports the peak potentials of the first and second reductions. As expected, the presence of -CF3 (1a and 1b) and –CN (1c) shifts first and second potentials toward more positive values, when compared to Mn(bpy)(CO)3Br. In a recent paper (Rawat et al., 2019) DFT calculations suggested that electron-withdrawing substituents like -CF3 stabilize the radical anion. Furthermore,
the formation of all Mn-Mn dimers was indicated as unfavorable. However, the electrochemical mechanism outlined above is commonly accepted, and the Mn dimer is strongly favored. We experimentally found that all complexes undergo a first and second chemically irreversible reductions. For example, 1a shows two chemical irreversible processes followed by the reoxidations of the pentacoordinated radical anion and that of the dimer (Figure S10), even at high scan rates (1 V/s, Figure S11), thus confirming the general mechanism. CV of 1g confirms that the strong electron-donating properties of dimethyl amino group result in more negative reduction potentials with respect to Mn(bpy)(CO)3Br. The push-pull system 1d, 1e and 1f display similar potential values to the unsubstituted bpy-complex. In some complexes decomposition processes occurring after reduction generate small peaks (i.e., for 1e and 1f E_p = −1.65 and −1.63 V vs. Ag/AgCl are observed, respectively). The very negative peak around −3V vs. Ag/AgCl, is commonly assigned to a ligand-centered reduction.

CVs of all rhenium complexes are reported in Figure 3. CV of the reference Re(bpy)(CO)3Cl under our experimental conditions is included for comparison. This complex exhibits a first reversible reduction, due to the formation of the radical anion, which is more stable than the analogous with manganese, thus resulting in electrochemical reversibility and no presence of the reoxidation peak of the dimer (Hawecker et al., 1984). The second reduction leads to the pentacoordinated anion. Intrinsically, rhenium complexes require slightly higher overpotentials with respect to the corresponding manganese ones. The first reversible reduction of Re(bpy)(CO)3Cl is located at E_1/2 = −1.35 V vs. Ag/AgCl, whereas the second chemically irreversible reduction is at E_p2 = −1.80 V vs. Ag/AgCl. Similarly to manganese, the new synthesized rhenium complexes show no significant difference in the electrochemical pathway under Ar when compared to the reference Re(bpy)(CO)3Cl (Table 2). Electron withdrawing groups (2a, 2b, and 2c) shift the reduction processes toward less negative values; electron donating groups like in 2g essentially merge the first and second reductions into a single peak. The push-pull system 2d, in analogy with the case of 1d, do not significantly alter the positioning of the reduction potentials. The third peak, around −2.5V, also for these complexes, is generally attributed to a bpy-centered reduction.

The reduction potentials estimated by DFT calculation are in excellent agreement with the experimental values obtained for the first reduction potential of all rhenium complexes (Figure 4). This confirms the reversible nature of the first reduction process, leading to stable radical anions characterized by a very small increase (about 0.05 Å) in the Re–Cl bond length (Figure S12). Conversely, in the case of manganese complexes, DFT calculations, which compute thermodynamic reduction potentials, are not suitable to estimate the irreversible electrochemical behavior of the compounds (Figure S13). Indeed, the high instability of the radical anion leads to the weakening of the Mn–Br bond, as clearly evidenced by the significant increase of the Mn–Br bond length in the anion structures (Figure S14). Our DFT calculations that include weak interactions, are in agreement with the chemically irreversibility of the first reduction and dimer production, even in the case of electron-withdrawing substituents. For example, the formation of the dimer [Mn(bpy)(CO)3]2⁺ from its radical anion is favored by 143.5 kJ/mol, and in the case of 1a is favored by 102.6 kJ/mol.

**Cyclic Voltammetry Under CO2**

The electrochemical behavior of manganese complexes under CO2 and with H2O (5%) is reported in Figure 5 for 1d to
CVs of 1 mM solutions of 1a−1g Mn complexes in MeCN/0.1 M TBAPF$_6$ at GCE, scan rate 200 mVs$^{-1}$ under Ar. CV of the reference fac-Mn(bpy)(CO)$_3$Br is in black.

1g and in Figure S15 for 1a, 1b, and 1c, (these complexes are catalytically inactive toward CO$_2$ reduction). CVs of 1a-1g under CO$_2$ with 5%v MeOH are included for comparison in Figure S15 too. All complexes do not show significant current increases switching from Ar to CO$_2$ atmosphere.

### Table 2: Reduction peak potentials [V] from the CV of Mn and Re complexes.

| Mn complexes | $E_{p1}$ | $E_{p2}$ |
|--------------|----------|----------|
| Mn(bpy)(CO)$_3$Br | −1.29 | −1.51 |
| 1a | −0.92 | −1.07 |
| 1b | −0.88 | −0.99 |
| 1c | −0.89 | −0.89 |
| 1d | −1.31 | −1.42 |
| 1e | −1.26 | −1.4 |
| 1f | −1.24 | −1.39 |
| 1g | −1.64 | −1.91 |

| Re complexes | $E_{1/2}$ (or $E_{p1}$) | $E_{p2}$ |
|--------------|---------------------|----------|
| Re(bpy)(CO)$_3$Cl | −1.35 | −1.80 |
| 2a | −0.92 | −1.45 |
| 2b | −0.83 | −1.57 |
| 2c | −0.77 | −1.85 |
| 2d | −1.29 | −1.77 |
| 2e | −1.31 | −1.68 |
| 2f | −1.37 | −1.66 |
| 2g | −1.82 (1$^{st}$ + 2$^{nd}$ peak) |

While this is expected for manganese complexes, 1g shows a current increase, though limited, even in absence of Brønsted acid addition.

The electrochemical behavior of rhenium complexes under CO$_2$ and with 5%v MeOH is reported in Figure 6 for 2d to 2g and in Figure S16 for 2a, 2b, and 2c. CVs of 2a-2g under CO$_2$ with 5%v H$_2$O are included for comparison in Figure S16 too. All complexes exhibit catalytic current switching from Ar to CO$_2$ atmosphere, even in absence of Brønsted acid, as expected for rhenium complexes.

### Controlled Potential Electrolysis Experiments of the Complexes

Bulk electrolysis experiments of all manganese and rhenium complexes series under CO$_2$ were performed upon setting the potential at values slightly negative to the second reductions, with and without external added Brønsted acids (water and methanol, 5%). A CO$_2$ flow of 50 mL min$^{-1}$ was kept constant during the experiments, gaseous products were determined by gas chromatography, and formate, if present, was assessed by NMR spectroscopy at the end of the experiments. Table 3 summarizes the results obtained during these CPE experiments. A general trend can be outlined: addition of water results in increased TONs for Mn complexes (Table 3 and Figure S17), differently from methanol, which drops them to lower values (Table S7). On the other hand, for the case of Re complexes, addition of methanol (Table 3 and Figure S18) seems to enhance the catalytic activity with respect to water, except for complex 2d, which is catalytically inactive, despite the promising current increase in presence of MeOH under CO$_2$. 

**TABLE 2 | Reduction peak potentials [V] from the CV of Mn and Re complexes.**
CONCLUSIONS

In summary, a systematic study of the effect of the electronic properties of the substituents on 2,2′-bipyridine Mn and Re complexes was conducted. Electron-withdrawing substituents shift the reduction potentials to more positive values, and eventually inhibit the catalytic activities of the corresponding Mn and Re complexes toward CO₂ reduction. In the case of electron-donating substituents the opposite trend is observed. These observations are in agreement with the induced electron density localized on the metal, strongly influencing the change in the reactivity with the weak electrophile CO₂. Increasing or decreasing the electron density on the metal should facilitate or prevent the formation of intermediate in which the CO₂ is coordinated to the metal. Another interesting effect in varying the electron properties of the substituents is the merging of the first and second reduction processes, observed in some Re and Mn complexes. A judicious selection of bpy substituents provides an alternative way to the use of bulky substituents to prevent dimer formation (Sampson et al., 2014), with the aim of transforming two 1e reduction into a single 2e reduction process (CO₂ to CO reduction requires 2e).

It is interesting to note that in spite of the presence of an OH group in 1f, no formic acid is detected, probably because in contrast with Mn carrying local proton sources (Franco et al., 2014, 2017) the hydroxyl group is located far from the metal center, thus the generation of the metal-hydride, commonly considered catalyst for formate production, is no longer entropically favored. It is interesting to note how CVs under CO₂ of the push-pull systems 1d and 2d show enhanced catalytic currents; during CPE the Mn derivative 1d displays the higher TON value, while the corresponding Re derivative 2d, albeit from CV appears to be a potentially highly active catalyst, undergoes decomposition. While Mn catalysts suffer from the presence of MeOH, they appear to work better in water, which seems to react promptly with reduced Mn. In fact, even if the reduction potential of 1g is rather negative (Table 2), no hydrogen is produced. This is in line with the high TON values observed for the CO₂ electrochemical reduction in pure water by Mn electrocatalysts supported on electrode surface (Walsh et al., 2015; Reuillard et al., 2017; Rotundo et al., 2019b). DFT calculations performed with dispersion correction agree with...
FIGURE 5 | CVs of 0.5 mM solutions of 1d–1g Mn complexes in MeCN/0.1 M TBAPF$_6$ at GCE, scan rate 200 mV s$^{-1}$ under Ar, under CO$_2$ and with H$_2$O (5%). CV in black is the electrolyte saturated with CO$_2$.

the experimental data for both Mn and Re complexes. The Mn–Br bonds computed for the Mn radical anions undergo a significant elongation, around 0.2 Å, which indicate non-negligible weakening of the Mn–Br bonds. The release of halogen is also very probably favored by the polar and coordinating solvent MeCN; indeed Br substitution by MeCN has been experimentally observed not only in the radical anion, but also in neutral Mn species (Franco et al., 2017). We demonstrated here how the appropriate choice of the electron properties of the ligands is of critical importance in the design of more effective bipyridine Mn and Re electrocatalysts for CO$_2$ reduction.

MATERIALS AND METHODS

General Considerations

CV and CPE experiments were performed using a Metrohm Autolab 302n potentiostat. CO and H$_2$ as CO$_2$ reduction products were detected and quantified by an Agilent 490 Micro GC. NMR spectra were recorded on a JEOL ECP 400 FT-NMR spectrometer ($^1$H operating frequency 400 MHz) or on a JEOL ECZR 600 FT-NMR spectrometer ($^1$H operating frequency 600 MHz) at 298 K. $^1$H and $^{13}$C chemical shifts are reported relative to TMS ($\delta = 0$) and referenced against solvent residual peaks. IR-ATR spectra were collected on a Fourier transform Equinox 55 (Bruker) spectrophotometer equipped with an ATR device; resolution was set at 2 cm$^{-1}$ for all spectra. A spectral range of 400–4,000 cm$^{-1}$ was scanned, using KBr as a beam splitter. GC-MS spectra were obtained on a mass selective detector Agilent 5,970 B operating at an ionizing voltage of 70 eV connected to a HP 5,890 GC equipped with a HP-1 MS capillary column (25 m length, 0.25 mm I.D., 0.33 µm film thickness. Elemental analyses (C, H, N) were performed on a Fisons Instruments EA-1108 CHNS-O Elemental Analyzer. ESI-MS spectra were recorded with a Thermo Advantage Max Spectrometer equipped with an ion trap analyzer and an ESI ion source.

Synthesis and Characterization of Ligands

Flasks and all equipment used for the generation and reaction of moisture-sensitive compounds were dried by electric heat gun under N$_2$. All commercially obtained reagents and solvents were used as received. Anhydrous DMF was purchased by Sigma Aldrich. Products were purified by preparative column chromatography on Macherey Nagel silica-gel for flash chromatography, 0.04–0.063 mm/230–400 mesh. Reactions were monitored by TLC using silica-gel on TLC-PET foils Fluka, 2–25 µm, layer thickness 0.2 mm, medium pore diameter
60 Å. hbppy (4-(2-hydroxy-phenyl)-6-phenyl-2,2'-bipyridine) was synthesized by using the Kröhnke (1976) reaction and subsequently coupling the pyridinium iodide salt (1) with the corresponding hydroxy chalcone (2). dpbpy (4,6-diphenyl-2,2'-bipyridine) was synthesized according to reported procedure (Franco et al., 2015).

**Synthesis of 4,4'-bis(trifluoromethyl)-2,2'-bipyridine:** the method of O’Donnell et al. (2016) was slightly modified to synthesize 4,4'-bis(trifluoromethyl)-2,2'-bipyridine first reported by Furue et al. (1992). In a degassed 20 mL screw cap vial a solution of 2-bromo-4-(trifluoromethyl)pyridine (500 mg, 2.2 mmol) in 7.5 mL of anhydrous DMF was poured and degassed for 5 min. Pd(OAc)$_2$ (25 mg, 0.11 mmol) was added later and the mixture degassed for additional 5 min. TBAI (815 mg, 2.2 mmol), anhydrous K$_2$CO$_3$ (460 mg, 3.3 mmol), and i-PrOH (0.35 mL, 4.4 mmol) were then added to the mixture, subsequently heated at 100°C for 20 h. The heating was suspended, and the reaction mixture was filtered through a pad of celite. The filtrate was diluted in 25 mL of DCM and was washed with deionized water (3 × 25 mL). The organic phase was collected, and the water layer further extracted with 10 mL of DCM. The collected organic phase was dried with anhydrous MgSO$_4$, filtered, and the solvent removed under reduced pressure to afford the crude solid that was purified by column chromatography on silica gel (eluent PE/AcOEt 10/1) to afford 220 mg of white solid as product. (Yield = 80%).

**Synthesis of [2,2'-bipyridine]-4,4'-dicarbonitrile:** the procedure reported by Losse et al. (2008) was followed with slight modifications. 4-Cyanopyridine (700 mg, 6.7 mmol) and 10% Pd/C (50 mg) were added into a 25 mL round-bottomed flask and five vacuum/nitrogen cycle were operated to minimize the amount of oxygen, then the flask was connected to a reflux condenser under nitrogen atmosphere. The mixture was heated to 230°C in a sand bath in order to reflux the 4-Cyanopyridine. After 24 h the mixture was cooled to room temperature, CHCl$_3$ (15 mL) was added and the black suspension filtered through a frit. Solvent was removed under reduced pressure from the pale-yellow solution obtained until the product started to crystallize. Pentane (15 mL) was then added, and the concentrated solution cooled at 4–6°C for 16 h. The precipitate was filtered on Buchner funnel and washed with cold EtOH and dried to yield 100 mg of the product as a yellow-orange solid. (Yield 15%).

**General procedures for synthesis of 2-substitued N,N-dimethylpyridin-4-amine derivatives.** The previously reported procedure of Cuperly et al. (2002) was adapted according to the used electrophile. 2-(dimethylamino)-ethanol (0.8 mL, 8.0 mmol) was added to a three-necked round bottomed flask and dissolved in hexane (10 mL) under a N$_2$ atmosphere. The
solution was cooled at −5°C, BuLi (2.5 M, 6.4 mL, 16.0 mmol) was added dropwise for 10 min and the resulting mixture stirred for 40 min at −5°C. 4-DMAP (488 mg, 4.0 mmol) was then added and stirring continued for additional 60 min at 0°C, then the reaction medium was cooled at −78°C and the solution of the appropriate electrophile (10.0 mmol) was added dropwise by mean of a dropping funnel with pressure balance. Once the addition of the electrophile was completed the temperature was allowed to raise to 0°C (1.5 h) and the reaction quenched with deionized water at this temperature.

Synthesis of N,N-dimethyl-2-(tributylstannyl)pyridin-4-amine. 2-(dimethylamino)ethanol (0.8 mL, 8.0 mmol) in hexane (10 mL), BuLi (2.5 M, 6.4 mL, 16.0 mmol), and 4-DMAP (0.488 g, 4.0 mmol) were reacted as previously described then a solution of Bu3SnCl (3.255 g, 10.0 mmol) in 15 mL of hexane was added dropwise for 20 min. The reaction was quenched with deionized water (15 mL) then the a dozen phase was extracted with DCM (2 × 20 mL) and AcOEt (2 × 20 mL). The collected organic phase was dried with anhydrous Na2SO4, filtered, and the solvent removed under reduced pressure to afford a crude orange oil, that was used without further purification in the cross-coupling reactions (NMR-calculated yield = 69%).

Synthesis of 2-iodo,N,N-dimethylpyridin-4-amine. 2-(dimethylamino)ethanol (1.6 mL, 16.0 mmol) in hexane (25 mL), BuLi (2.5 M, 12.8 mL, 32.0 mmol), and 4-DMAP (0.980 g, 8.0 mmol) were reacted as previously described then a solution of resublimed I2 (5.080 g, 20.0 mmol) in 50 mL of freshly distilled Et2O was added dropwise for 35 min. The reaction was quenched with a saturated solution of Na2S2O3 (25 mL) and stirred for additional 20 min at 0°C. The organic phase was separated and washed again with Na2S2O3 solution (2 × 15 mL) and brine (2 × 15 mL). The collected organic phase was dried with anhydrous Na2SO4, filtered, and the solvent removed under reduced pressure to afford a crude brown solid that was purified by column chromatography on silica gel (eluents: PE/AcOEt 5/5) to afford 1.683 g of white solid as product (Yield = 86%).

General Procedure for Stille cross-coupling of N,N-dimethyl-2-(tributylstannyl)pyridin-4-amine with 2-halopyridine. In a degassed 50 mL screw cap vial freshly distilled toluene (25 mL) was added and degassed for 10 min then Pd(OAc)2 (22 mg, 0.1 mmol) and PPh3 (52 mg, 0.2 mmol) were added in one portion. The resulting mixture was stirred and degassed until toning of the solution to red. Subsequently, N,N-dimethyl-2-(tributylstannyl)pyridin-4-amine (370 mg, 0.9 mmol), LiI (40 mg, 0.3 mmol), and CuI (57 mg, 0.3 mmol) were added to the mixture, then degassed for additional 5 min. Finally, the appropriate 2-halopyridine (1.1 mmol) was poured in the reaction medium that was kept under N2 atmosphere and heated to reflux for 16 h. After being cooled at room temperature, the resulting mixture was diluted with EtOAc (30 mL) and washed with a NH4OH solution (10 M) until the water layer did not turn blue any more, indicating that all the copper had been extracted. The collected organic phase was filtered on a pad of celite, diluted with DCM (30 mL) and dried with anhydrous Na2SO4. Then the solvents were removed under reduced pressure to afford the crude solid subsequently purified to yield the desired product.

Synthesis of N4,N4',N4'-tetramethyl-[2,2'-bipyridine]-4,4'-diamine: Following the general procedure, N,N-dimethyl-2-(tributylstannyl)pyridin-4-amine (370 mg, 0.9 mmol) and 2-iodo-N,N-dimethylpyridin-4-amine (270 mg, 1.1 mmol) in 25 mL of freshly distilled toluene were reacted to afford a crude yellowish solid that was suspended in 25 mL of Et2O and vigorously stirred for 30 min. The desired product was separated by filtration to yield 115 mg of a brown powder (Yield = 53%).

Synthesis of N,N-dimethyl-4'-((trifluoromethyl)-[2,2'-bipyridin]-4-amine: following the general procedure, N,N-dimethyl-2-(tributylstannyl)pyridin-4-amine (370 mg, 0.9 mmol) and commercially available 2-bromo-4-((trifluoromethyl)pyridine (250 mg, 1.1 mmol) in 25 mL of freshly distilled toluene were reacted to afford a crude brown solid that was purified by column chromatography on silica...
gel (eluent: MeOH/DCM 5/95) to yield 169 mg of the desired product as a brown powder (Yield = 70%).

**Synthesis of N-(2-pyridylacetyl)-pyridinium iodide (1):** in a three-necked flask 82.5 mmol of 2-acetylpyridine and 90.5 mmol of iodine are dissolved in 100 mL of pyridine. The solution is refluxed for 3 h. The shiny black precipitate is then filtered and washed with diethyl ether (20 mL). The solid is recrystallized in hot ethanol: fine-scaled golden crystals are obtained (overall yield: 54%).

**Synthesis of (E)-3-(2-hydroxyphenyl)-1-phenyl-2-propen-1-one (2):** is synthesized by Claisen-Schmidt condensation of salicyaldehyde (50 mmol) with acetoephone (40 mmol) dissolved in 50 mL of ethanol (Tatsuzaki et al., 2006; Yin et al., 2012). The solution is vigorously stirred and 10 mL of KOH 40% (w/w) is dropwise added. The mixture is then heated to 60°C for 2–4 h until the disappearance of the starting reagents, monitored by TLC (EtAc: hexane 1:4). Once the reaction is completed, the suspension is poured into cold distilled water and acidified with HCl 2 M (final pH 2–3). The resulting precipitate is collected and washed once with water and then with cyclohexane (yield: 70%).

**Synthesis of 4-(2-hydroxy-phenyl)-6-phenyl-2,2’-bipyridine** (hpbpy): precursors 1 (4.8 mmol) and 2 (4.8 mmol) are added to a three-necked flask with an excess of ammonium acetate (48 mmol) and glacial acetic acid (4 mL). The solution is refluxed for 3 h. The glacial acetic acid is removed, and the residue dissolved in methanol. The yellowish product cristallizes on cooling.

**Synthesis of 4,6-diphenyl-2,2’-bipyridine** (dpbpy): is already reported in literature; NMR data are in agreement with previously published data (Cave and Raston, 2001).

**Elemental Analysis of the Complexes**

The samples for microanalyses were dried in vacuum to constant weight (20°C, ca. 0.1 Torr). Elemental analysis (C, H, N) was performed in-house with a Fisons Instruments 1108 CHNS-O Elemental Analyzer.

| Complex | Anal. Calcd. (%) | Found (%) |
|---------|-----------------|-----------|
| 1       | C, 35.55; H, 1.25; N, 5.45 | C, 35.55; H, 1.09; N, 5.57 |
| 2       | C, 30.20; H, 1.12; N, 4.35 | C, 30.13; H, 1.01; N, 4.69 |
| 3       | C, 48.90; H, 2.63; N, 4.45 | C, 48.90; H, 2.63; N, 4.45 |

**Single-Crystal X-Ray Diffraction**

The single-crystal data were collected with a Gemini R Ultra diffractometer with graphite-monochromated Mo-Kα radiation (λ = 0.71073) by the o-scan method. The cell parameters were retrieved with the CrysAlisPro (Agilent, 2015) software, and the same program was used to perform data reduction with corrections for Lorentz and polarizing effects. Scaling and absorption corrections were applied through the CrysAlisPro1 multiscan technique. The structures of complex 2e (both structure A and B) were solved with direct methods, while in the case of 2f a meaningful initial guess for electron density was obtained only with Patterson Function by using SHELXS-14 (Sheldrick, 2008, 2015). All the structures were refined with full-matrix least-squares techniques on F² with SHELXL-14 (Macrae et al., 2006) using the program Olex2 (Dolomanov et al., 2009). All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were calculated and riding on the corresponding bonded atoms. The graphic of the crystal structures was generated using Mercury 3.9 (Macrae et al., 2006). CCDC codes 1891407–1891409 contain the supplementary crystallographic data for 2e (structure A), 2e (structure B) and 2f. These data can be obtained free of charge via https://www.ccdc.cam.ac.uk/conts/retrieving.html, or from the Cambridge Crystallographic Data Center, 12
CV and CPE Experiments

Acetonitrile used for the experiments was freshly distilled over calcium hydride and purged with Ar before use. 0.5–1 mM solutions of the complexes were prepared with tetrabutylammonium hexafluorophosphate (TBAPF₆, Sigma-Aldrich, 98%) as supporting electrolyte (0.1 M). A single-compartment cell was employed for CV measurements, equipped with working a glassy carbon electrode (GCE, Ø = 1 mm), alongside a Pt counter electrode and a Ag/AgCl (KCl 3 M) reference electrode. The Ar- and CO₂-saturated conditions were achieved by purging gases for 5 min before each potential sweep. A double-compartment H-type cell was used for CPE measurements, thus allowing to separate through a glass frit the anodic compartment from the cathodic one (a Pt wire is placed as counter electrode). A glassy carbon rod is employed as working electrode jointly with the Ag/AgCl reference electrode. A controlled and constant flow of CO₂ (50 mL min⁻¹) was maintained during the CPE measurements by means of a Smart Trak 100 (Sierra) flow controller. Under these experimental conditions, the redox couple (Fc⁺/Fc) is located at E₁/₂ = 0.35 V.

Quantitative Analysis of CO₂ Reduction Products

µGC measurements were used to detect CO and H₂. Two modules equipped with CP-Molsieve 5 Å columns were kept at 105°C and at a pressure of 30 and 28 psi, with a thermal conductivity detector. The carrier gases were Ar for H₂ and He for CO detection, respectively. The backflush vent option time was set to 7 s. The gas inside the measurement cell was sampled for 30 s every 3 min to fill the Micro GC 10 µL sample loop, and eventually 500 nL was injected into the column for the analyses. Instrument calibration was carried out measuring two different certified standards of CO and H₂ in Ar matrix (Rivoira). Formate production was assessed by NMR spectroscopy.

Computational Details

All the calculations were performed by the Gaussian 16 Revision B.01 (G16) program package (Frisch et al., 2016), employing density functional theory (DFT). Calculations were run using the Becke three-parameter hybrid functional (Becke, 1993), and the Lee–Yang–Parr gradient-corrected correlation functional (B3LYP) (Lee et al., 1988). Dispersion effects were added as semiempirical corrections with Becke–Johnson damping approach (GD3BJ) (Grimme et al., 2010, 2011). The solvent effect was included using the conductor-like polarizable continuum model (CPCM) with acetonitrile as solvent (Miertus et al., 1981). The def2TZVP basis set and effective core potential were used for the Mn, Br, and Cl atoms and the def2-SVP basis set was used for all the other atoms (Weigend and Ahlrichs, 2005). Unrestricted open-shell calculations were performed on the radical anions. Geometry optimizations were carried out without any symmetry constraints. The nature of the stationary points in the potential energy hypersurface was characterized by using harmonic vibrational frequency calculations. No imaginary frequencies were found, thus indicating we had located the minima on the potential-energy surfaces. Molecular-graphics images were produced by using the UCSF Chimera package from the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco (Pettersen et al., 2004).

AUTHOR CONTRIBUTIONS

RG and CN as corresponding authors wrote and revised the manuscript. EP performed Crystal X-ray structures. EA, AD, and PQ made the synthesis of the ligands. LR, RR, and LN synthesized the organometallic catalysts and performed the electrochemical and GC measurements. CG and CN made the DFT calculations.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2019.00417/full#supplementary-material

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