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Pre-equilibrium reactions involving pendent relays

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Homogeneous earth abundant transition-metal electrocatalysts capable of carbon dioxide (CO₂) reduction to generate value-added chemical products are a possible strategy to minimize rising anthropogenic CO₂ emissions. Previously, it was determined that Cr-centered bipyridine-based N₂O₂ complexes for CO₂ reduction are kinetically limited by a proton-transfer step during C-OH bond cleavage. Therefore, it was hypothesized that the inclusion of pendent relay groups in the secondary coordination sphere of these molecular catalysts could increase their catalytic activity. Here, it is shown that the introduction of a pendent methoxy group favorably impacts a pre-equilibrium protonation prior to the catalytic resting state, resulting in a significant increase in catalytic activity without a loss of product selectivity for generating carbon monoxide (CO) from CO₂. Interestingly, combining the pendent methoxy group with a cationic acid causes a positive shift of the catalytic reduction potential of the system, while maintaining increased activity and quantitative selectivity. This work suggests that tuning the secondary coordination sphere with respect to cationic proton sources can result in activity improvements by modifying the kinetic and thermodynamic aspects of proton transfer in the catalytic cycle.

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Introduction

The industrial revolution marked the start of a dramatic increase in the amount of accumulated atmospheric carbon dioxide (CO₂), a leading contributor to the global climate crisis. As of 2021 the amount of atmospheric CO₂ is approximately 1.4 trillion tons, compared to approximately 45 billion tons in 1900.2 Therefore, there is significant interest in mitigating the impact of increased solar energy retention caused by elevated levels of atmospheric CO₂. Electrochemically reducing CO2 to value-added chemical products through the use of renewable energy sources, such as wind or solar, is one way to lessen the concentration of CO₂ in the atmosphere.³ The reduction of CO2 can be accomplished through protoncoupled multi-electron pathways, but the majority of molecular electrocatalysts reduce CO₂ via a two-electron, two-proton (2e⁻/ 2H⁺) mechanism, producing either carbon monoxide (CO) with water (H₂O) as a co-product or formic acid (HCOOH).⁴ Both carbon-containing products are relevant to industrial applications: the Fischer-Tropsch process uses CO and H2 to

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produce fuels and commodity chemicals⁵ while formic acid is an energy dense carrier for fuel cells, as well as a useful chemical synthon.⁶

Although the CO2 reduction reaction (CO2RR) can be mediated by homogeneous complexes based on expensive late transition metals (Pd,7 Ru,8 and Re9), a more sustainable approach would be to use earth-abundant transition metals to convert CO₂ into these useful products for chemical feedstocks or fuels. 10 In 1984, a [Ni^{II}(cyclam)]²⁺ catalyst for the selective reduction of CO₂ to CO on a mercury cathode was reported by Beley et al.,11 following up on early work from Fisher and Eisenberg in 1980 on less-selective reaction conditions. ¹² Since then, there have been other examples of Earth-abundant transition metal catalysts for the CO2RR, most notably, manganese (Mn) carbonyl bipyridyls¹³ and iron (Fe) porphyrins.¹⁴ The most active molecular catalyst reported to date is the iron porphyrin, Fe-o-TMA (iron(III) 5,10,15,20-tetra-(o-N,N,N-trimethylanilium)-porphyrin), reported by Azcarate et al., which has a turnover frequency (TOF) of $10^6 \text{ s}^{-1}.^{15,16}$

Although for decades chromium (Cr) was thought to be nonprivileged for the $\mathrm{CO}_2\mathrm{RR},^{17,18}$ more recently four examples of molecular Cr catalysts for the selective reduction of CO_2 to CO have been reported. $^{10,19-22}$ The first selective homogeneous Cr-based catalyst was $\mathrm{Cr}(^{tbu}\mathrm{dhbpy})\mathrm{Cl}(\mathrm{H}_2\mathrm{O})$, where the ligand precursor, $(^{tbu}\mathrm{dhbpy}(\mathrm{H})_2)$ is 6,6'-di(3,5-di-tert-butyl-2-hydroxybenzene)-2,2'-bipyridine. 10,19,20 The catalytic cycle for the CO_2 reduction reaction mediated by $\mathrm{Cr}(^{tbu}\mathrm{dhbpy})\mathrm{Cl}(\mathrm{H}_2\mathrm{O})$ proposed

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$$-CO, -S$$

$$-CO,$$

Fig. 1 Proposed catalytic mechanism for electrocatalytic CO_2 reduction mediated by $Cr(^{bu}dhbpy)Cl(H_2O)$.

by Hooe et al. under protic conditions is depicted in Fig. 1. 19,23 After an initial overall two-electron reduction of the Cr complex with chloride loss to form the four-coordinate monoanionic Cr(II)(bpy -) active species, i, CO₂ binds to the Cr center forming [Cr-CO₂]⁻, ii. The CO₂ adduct is subsequently protonated to form a neutral hydroxycarbonyl, iii. A second reduction with a standard potential more positive than $E_{\rm cat}^{\circ}$ monoanionic hydroxycarbonyl species [Cr-CO₂H]⁻, iv. Another proton transfer reaction cleaves the C-OH bond and releases water, leaving CO bound to the Cr center, v. The release of CO from this neutral species is expected to be rapid and generate a neutral four-coordinate species, vi, from which one-electron reduction at E_{cat}° closes the catalytic cycle. Density Functional Theory (DFT) calculations suggest that the cleavage of the C-OH bond to release water is the rate-determining step (RDS),²³ consistent with the experimental mechanistic data. During electrolysis, this Crbased complex produced CO with a FE of 96 ± 8% from CO₂ in the presence of 0.6 M phenol (PhOH).¹⁹

Pendent proton (or proton donor) relays are commonly used to facilitate proton transfer by means of hydrogen bonding and proximity to the active site during electrochemical reactions. Pendent relays incorporated into the secondary coordination sphere have been studied extensively for reductive small molecule conversion in the context of the oxygen reduction reaction, Secondary and the CO₂RR. Since the kinetic enhancement originates from the overall balance of the equilibria involving proton transfer from solution to the active site via the relay, secondary it is advantageous to have acid-base pairs with fast

exchange reactions (e.g., O or N containing).²⁴ For instance, in 2017, Ngo et al. described how the introduction of methoxy (-OMe) groups into a 2,2'-bipyridine-based ligand appended to a Mn(CO)₃Br core led to an increase in CO₂RR catalysis.⁵⁸ For this Mn-based system, the pendent Lewis base is able to form noncovalent hydrogen-bonding interactions with the C-OH intermediate, resulting in an overall stabilization of the C-OH bond cleavage step by acting as a proton donor relay. 58 Along with its increased catalytic activity, the system operated at lower overpotentials, as compared to the parent non-substituted catalyst, and controlled potential electrolysis (CPE) studies of the complex confirmed high product selectivity for CO in the presence of nonaqueous Brønsted acid proton sources. 58 In the context of the CO2RR, we have similarly examined a molecular Fe-based CO2RR catalyst with methoxy-based pendent relays incorporated into an analogous coordination environment to the Cr-based compound described above. 45 In this Fe system, the addition of the -OMe group increases the availability of the proton donor through non-covalent interactions, resulting in an increase in catalytic activity and improved selectivity for formate (HCO₂⁻).⁴⁵

Based on the proposed rate-determining C-OH bond cleavage step for Cr-based complexes during electrocatalytic CO_2 reduction, it was hypothesized that the addition of pendent relays to the ligand framework would have a beneficial effect on the mechanism of proton transfer, without resulting in significant changes to the electronic structure of the Cr active site (changes in E_{cat}°). Here, the synthesis, characterization, and electrochemical behavior of two new Cr-based complexes with a series of proton sources under unbuffered (acid only) and buffered conditions (acid and conjugate base added) under electrocatalytic conditions for the CO_2RR is reported. Notably, significant differences in activity and operating potential are observed when comparing a control complex, $Cr(^{\text{p-tbu}}\text{dhbpy})Cl(H_2O)$ (1) (Fig. 2) with a compound containing pendent –OMe groups, $Cr(^{\text{nPr}}\text{dhbpy})Cl(H_2O)$ (2) (Fig. 2). Using cyclic voltam-

tbu
$$Cr(^{p-tbu}dhbpy)Cl(H_2O)$$
(1)

 $Cr(^{p-tbu}dhbpy)Cl(H_2O)$
(1)

 $Cr(^{nPr}dhbpy)Cl(H_2O)$
(2)

Fig. 2 Structures of $Cr(^{p-tbu}dhbpy)Cl(H_2O)$ (1), and $Cr(^{nPr}dhbpy)Cl(H_2O)$ (2), where S is a solvent adduct of water or DMF.

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metry (CV) and CPE, the effects of the pendent relay on catalysis are elucidated, specifically the shift to a pre-equilibrium catalytic response that causes both a kinetic effect (higher TOF in CPE studies) and a thermodynamic effect (positive shift in catalyst operating potential) on CO2 reduction, without loss of quantitative product selectivity for CO. The pre-equilibrium approximation for catalysis applies when there is a fast initial step under equilibrium control prior to the rate-determining step, while the steady state approximation can be used to describe catalytic responses where the rate of intermediate formation is matched by the rate of intermediate consumption.

Results

Synthesis and characterization

The ligand precursor complex 6,6'-di(5-tert-butyl-2-hydroxybenzene)-2,2'-bipyridine, (p-tbu dhbpy(H)2) was synthesized as previously reported.⁶⁰ The synthesis of the 6,6'-di(3-methoxy-5-npropyl-2-hydroxybenzene)-2,2'-bipyridine, (nPrdhbpy(H)2) ligand was completed via a modification of our previously reported procedure (see ESI† for details).²⁸ Metalation of $(^{p-tbu}dhbpy(H)_2)$ and $(^{nPr}dhbpy(H)_2)$ to generate $Cr(^{p-tbu}dhbpy)$ Cl(H2O) (1) and Cr(nPrdhbpy)Cl(H2O) (2), respectively, was achieved by refluxing a solution of ligand and Cr(II) chloride metal salt under inert atmosphere for 24 hours, analogous to previously reported procedures (see the ESI†).19 Purified products were characterized by UV-vis (Fig. S2-S4†), NMR (Tables S2 and S3†), and microanalysis. Crystals of 1 suitable for XRD studies were grown from a concentrated solution of 1 in N,Ndimethylformamide (DMF) at cold temperatures (Fig. S1†).

Electrochemistry

To probe the electrochemical behavior of these complexes, cyclic voltammetry (CV) was performed on 1 and 2 in DMF with 0.1 M tetrabutylammonium hexafluorophosphate (TBAPF₆) as the supporting electrolyte. Under argon (Ar) and CO₂ saturation conditions, three redox features were observed for 1 at $E_{1/2}$ = -1.59, $E_p = -1.75$, and $E_{1/2} = -1.95$ V *versus* ferrocenium/ferrocene (Fc⁺/Fc) (Fig. 3). The first two redox features of 1 coalesce upon the addition of excess chloride (Fig. S5 and S6†), consistent with the features being chemically related to each other by chloride loss. It is proposed that the more positive redox feature at $E_{1/2}$ = -1.59 V vs. Fc⁺/Fc represents a solvento species generated by an equilibrium chloride displacement reaction, while the more negative at $E_p = -1.75 \text{ V } \text{vs. Fc}^+/\text{Fc}$ corresponds to the neutral chloro adduct based on precedent. 19,20

For 1, under Ar and CO₂ saturation conditions, there are minimal changes observed in the CV response (Fig. 3). However, under both Ar and CO2 saturation, the addition of PhOH results in the appearance of a feature at a potential positive of the first reduction feature in the forward trace with E_p = -1.51 V vs. Fc⁺/Fc (Fig. 3). To probe the nature of this prewave feature observed for 1, CV responses with an excess of tetrabutylammonium chloride (TBACl) were conducted. In the presence of an excess of TBACl under Ar and CO2 saturation, a shift in the first reduction feature from $E_p = -1.59$ to -1.63 V vs. Fc⁺/Fc was observed, along with the disappearance of the second redox feature at $E_p = -1.75 \text{ V}$ vs. Fc⁺/Fc (Fig. S6 and S7†). The combination of excess TBACl and PhOH under Ar and CO₂ saturation conditions suppressed the appearance of the prewave observed in the presence of PhOH only (Fig. S8†), although it still could be observed at high scan rates (Fig. S6 and S7†). Similar to the addition of TBACl by itself, a shift to more negative reduction potentials for the original first reduction feature and disappearance of the original second redox feature was observed with the combination of TBACl and PhOH (Fig. S8†). Overall, these results agree with previously reported results19,23 and are consistent with the proposal that the two reduction features at $E_{1/2} = -1.59$ and $E_p = -1.75$ V vs. Fc⁺/Fc are related to an equilibrium chloride displacement reaction for complex **1**. The pre-wave feature at $E_p = -1.51 \text{ V}$ vs.

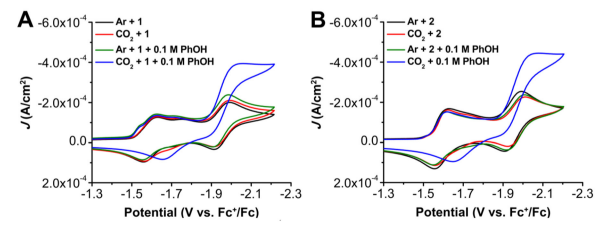


Fig. 3 (A) Comparison CVs of $Cr(^{p-tbu}dhbpy)Cl(H_2O)$ 1 under Ar and CO_2 saturation conditions with and without 0.1 M PhOH. (B) Comparison CVs of Cr(nPrdhbpy)Cl(H₂O) 2 under Ar and CO₂ saturation conditions with and without 0.1 M PhOH. Conditions: 1.0 mM catalyst, 0.1 M TBAPF₆/DMF; glassy carbon disc working electrode, glassy carbon rod counter electrode, Aq/AqCl pseudoreference electrode; referenced to Fc⁺/Fc internal standard; 100 mV s⁻¹ scan rate.

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Fc⁺/Fc can be ascribed to reversible and weak adsorption to the electrode, based on its shape and the rinse tests conducted under electrolysis conditions described below. The enhancement of this pre-wave with added phenol suggests that the proton donor could assist with chloride loss and therefore alter electrode adsorption.

Variable scan rate studies under both Ar (Fig. S6†) and CO₂ (Fig. S7†) were subsequently examined and the diffusion-controlled saturation response at the reversible third reduction feature ($E_{1/2} = -1.95 \text{ V } \nu s. \text{ Fe}^+/\text{Fc}$) is representative of a homogeneous electrochemical response under both conditions (Fig. S6 and S7†). In the presence of PhOH (p K_a = 18.8 in DMF)⁶¹ as a proton source under CO₂ saturation, a current increase indicative of a catalytic response for the CO2RR was observed to originate from the redox wave at $E_{1/2} = -1.95 \text{ V} \text{ vs.}$ Fc⁺/Fc (Fig. 3). Mechanistic analysis was attempted *via* variable concentration studies via a log-log plots in regions where catalytic current was sensitive to substrate concentration.⁶² These studies suggest a mixed-order concentration dependence of 1 (Fig. S9†), first-order dependence of PhOH (Fig. S10†), and mixed-order dependence of CO2 (Fig. S11†) on the observed electrocatalytic current. Variable scan rate studies establish that 1 is in the appropriate kinetically limited regime (Fig. S13†), suggesting that pre-equilibrium processes are being observed in the concentration dependence of the electrocatalytic response.

Analogous CV experiments for 2 under Ar and CO2 saturation similarly show three reduction features at $E_{1/2}$ = -1.59, $E_{\rm p}$ = -1.72, and $E_{1/2}$ = -1.95 V vs. Fc⁺/Fc (Fig. 3). The first two redox features of 2 coalesce with the addition of chloride (Fig. S33 and S34†), consistent with the features being chemically related to each other; as previously assigned, the more positive redox feature represents a solvento species generated by an equilibrium chloride displacement reaction while the more negative corresponds to the neutral chloro adduct. 19,20 Unlike complex 1, no adsorption pre-feature is observed either at high-scan rates or upon the addition of added proton donor to complex 2 (Fig. 3B), implying an improved solubility. As was

the case with 1, the addition of PhOH under CO2 saturation results in a large increase in current and loss of reversibility at the most negative redox feature $(E_{1/2} = -1.95 \text{ V } \text{vs. Fc}^+/\text{Fc})$. Variable concentration studies show that the current corresponding to electrochemical CO2 reduction mediated by complex 2 has a first-order concentration dependence with respect to complex 2 (Fig. S36†) and mixed-order dependences on CO₂ (Fig. S37†) and PhOH (Fig. S38†). The relative current densities achieved for 1 and 2 at 0.6 M PhOH under CO2 saturation was approximately the same, but less than for the prereported bpy-based catalyst under comparable conditions. 10,19-21 Unlike complex 1, 2 also exhibited a 10 mV shift to more positive $E_{\text{cat/2}}$ as PhOH concentration increased (Fig. S38†). In the absence of CO₂, this shift was not observed, indicating that this shift relates to a modification of either preequilibrium CO2 binding or protonation of a CO2-containing intermediate. A comparable shift was not observed for 1.

Next, a weaker proton source, trifluoroethanol (TFE) (estimated p $K_a = 35.4$ in MeCN, ⁶³ scaled to ~24.0 in DMF⁶⁰) was tested to assess the role of the pendent relay. Similar to PhOH, a large increase in current density and loss of reversibility is observed at the third reduction of 1 and 2 with 0.6 M TFE. For 1 there was again no shift in $E_{\text{cat/2}}$ with TFE, however, a shift to more positive potentials by 30 mV was observed for the $E_{\text{cat/2}}$ of 2. Variable concentration studies with 1 and 2 show that the electrochemical reaction under these conditions has a firstorder dependence on catalyst and CO2 concentration, as well as TFE for both complexes (Fig. S14-16 and S41-43†). Since these results do not appear to scale with the reported pK_a when compared to PhOH, it can be speculated that solvation and homoconjugation effects for TFE impose slightly different constraints on the reaction environment. Finally, the more acidic and cationic acid triethylammonium hexafluorophosphate (TEAHPF₆) ($pK_a = 9.25$ in DMF)⁶⁴ was examined. As was the case with PhOH, a large increase in current density accompanied by a loss of reversibility is observed at the third reduction feature of 1 under CO₂ saturation and in the presence of 20 mM TEAHPF₆ (Fig. 4). Variable concentration

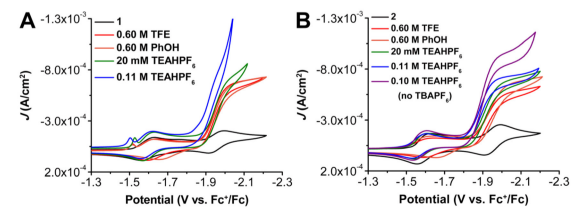


Fig. 4 Comparison CVs of (A) Cr(P-tbu dhbpy)Cl(H₂O) 1 and (B) Cr(P-tbu dhbpy)Cl(H₂O) 2 under CO₂ saturation conditions with different proton sources. Conditions: 1.0 mM catalyst, 0.1 M TBAPF₆/DMF unless otherwise noted; glassy carbon disc working electrode, glassy carbon rod counter electrode, Ag/AgCl pseudoreference electrode; referenced to Fc^+/Fc internal standard; 100 mV s⁻¹ scan rate.

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studies show that the electrochemical reaction under these conditions with 1 present has a first-order concentration dependence with respect to catalyst (Fig. S19†), however, mixed-order kinetics are again observed for TEAHPF₆ (Fig. S20†), and the catalytic waveform with CO2 shows evidence of an overlapping heterogeneous interaction between the proton donor and the electrode at low CO2 concentrations (Fig. S21†). It is worth emphasizing that increasing to 0.1 M concentrations of TEAHPF6 also results in significant changes to the waveform, suggestive of a heterogeneous interaction between the proton donor and the electrode (Fig. 4 and S26†). Consistent with this, comparing the i_{cat}/i_p ratio to the inverse square root of scan rate showed a linear response that deviated from intercept, indicating competing reactivity with the homogeneous process (Fig. S24†). Complex 1 was found to still exhibit an irreversible redox response consistent with the CO₂RR under buffered conditions (equivalent molar addition of conjugate base triethylamine (TEA) to solutions containing TEAH⁺), although the observed current density diminished significantly (Fig. S22†).

Under unbuffered conditions with 20 mM TEAHPF6 and complex 2, there is a slight increase in catalytic current density, with no waveform changes associated with an interaction between the proton donor and the electrode. Further, complex 2 experiences a 30 mV shift to more positive potentials, $E_{\text{cat/2}} = -1.92 \text{ V } \text{vs. Fc}^+/\text{Fc}$ (Fig. 4). It is proposed that the increased shift to more positive potentials for 2 in comparison to 1 is likely the result of the pendent methoxy groups interacting with the ammonium-based proton donor. Variable concentration studies show that the electrochemical reaction shows a mixed-order concentration dependence with respect to catalyst (Fig. S46†), TEAHPF₆ (Fig. S47†), and CO₂ (Fig. S48†); the concentration of the TEAHPF6 proton donor is fixed at 20 mM when the other reaction components are varied. Unlike 1, at 0.1 M TEAHPF₆ concentrations complex 2 remained catalytic for the CO₂RR ($E_{\text{cat/2}} = -1.89 \text{ V vs. Fc}^+/\text{Fc}$) without being overwhelmed by a heterogeneous electrode response. Interestingly, variable concentration studies conducted in the presence of the higher fixed concentration of 0.1 M TEAHPF₆ exhibited firstorder dependence with respect to catalyst (Fig. S49†) but the reaction order dependence on CO2 still showed a mixed-order response (Fig. S50†). Like 1, complex 2 was found to experience a slight decrease in catalytic activity when under buffered conditions (Fig. S51†). For complex 2 under unbuffered conditions, increasing the concentration of TEAHPF6 resulted again in a corresponding shift of $E_{\text{cat/2}}$ to more positive potentials; similar titration studies under these conditions did not show a shift of $E_{\text{cat/2}}$ for 1. Interestingly, measurements under buffered conditions did not reflect as significant a suppression of catalysis for 2 as was observed for 1.

The acid TEAHPF₆ is unique in that it has a relatively low saturation concentration in these CV systems (20 mM compared to 0.6 M PhOH or 1.1 M TFE with 2 in DMF) and has the potential to act as both an electrolyte and as an acid. CV studies were conducted on 1 in DMF with 0.1 M TEAHPF6 serving the dual roles of proton donor and supporting electrolyte. Under Ar saturation conditions, 1 has a redox feature that appears at $E_{1/2} = -1.59 \text{ V} \text{ vs. Fc}^+/\text{Fc}$ followed by a sharp increase in current that suggests the hydrogen evolution reaction (HER) is occurring, possibly originating from the electrode surface (Fig. S25†).65,66 Cross-tracing is also observed, implying either adsorption of homogeneous intermediates to the electrode surface or a slow chemical step leading to the accumulation of reactive species in the reaction-diffusion layer. 66 Under CO2 saturation conditions, two redox features appear at $E_{1/2} = -1.59$ V and $E_p = -1.88$ V vs. Fc⁺/Fc for 1, but the second feature is very slight and is quickly overcome by a large increase consistent with an HER response (Fig. S26†).65 Analogous CV experiments for 2 in DMF with 0.1 M TEAHPF6 as both proton donor and supporting electrolyte under Ar exhibited two redox features at $E_{1/2} = -1.57$ V and $E_p = -1.95$ V vs. Fc⁺/Fc (Fig. S26†). Under CO₂ saturation a large increase in current with a loss of reversibility is observed at the more negative redox feature, with $E_{\text{cat/2}} = -1.90 \text{ V} \text{ vs. Fc}^+/\text{Fc.}$ The relative current density of this system with 2 and 0.1 M TEAHPF6 as electrolyte and proton donor is also decreased compared to the system which also includes TBAPF₆ electrolyte.

Finally, CPE was performed to elucidate differences in product selectivity and assess TOF under preparative conditions for the different acids. CPE experiments for 1 with PhOH present under CO₂ saturation at an applied potential of -2.10 V vs. Fc⁺/Fc showed CO production with a Faradaic efficiency (FE_{CO}) of 85 ± 5% accompanied by minor H₂ production (Table 1), with a turnover frequency (TOF) of 0.29 s^{-1} . CPE experiments for 2 under comparable conditions demonstrated quantitative efficiency for CO (99 \pm 3%) and a slightly

Table 1 Results of CPE experiments under CO₂ saturation conditions

Conditions	Potential (V vs. Fc ⁺ /Fc)	FE_{CO} (%)	FE_{H_2} (%)	$TOF_{CPE}\left(s^{-1}\right)$	η (V)	Turnovers of CO w.r.t [1] or [2]
$1 + \text{PhOH}^{a,f}$	-2.10	85 ± 5	5 ± 0	0.29	0.09	2.14
$1 + \text{TFE}^{b,f}$	-2.10	103 ± 3	n/a	1.03	_	3.09
1 + TEAHPF ₆ c,f	-2.10	83 ± 1	17 ± 0	0.87	0.70	3.50
$2 + PhOH^{b,f}$	-2.10	99 ± 3	n/a	0.93	0.06	3.08
$2 + \text{TFE}^{b,f}$	-2.10	105 ± 12	n/a	1.50	_	2.56
$2 + \text{TEAHPF}_6^{d,f}$	-2.10	104 ± 4	n/a	13.70	0.66	4.61
$2 + \text{TEAHPF}_{6}^{e,g}$	-2.05	103 ± 5	n/a	5.47	0.66	5.05

^a 0.75 mM catalyst and 1.5 M acid. ^b 0.5 mM catalyst and 1.0 M acid. ^c 0.5 mM catalyst and 20 mM acid. ^d 0.4 mM catalyst and 16 mM acid. ^e 0.5 mM catalyst and 0.10 M acid. ^f 0.1 M TBAPF₆ present as electrolyte. ^g No electrolyte present.

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improved TOF of 0.93 s⁻¹. 10,21 CPE experiments with 1 where TFE was the proton donor produced CO with an efficiency of $103 \pm 3\%$ with a TOF of 1.03 s⁻¹. Complex 2 also showed quantitative efficiency of 105 \pm 12% and a TOF of 1.50 s⁻¹ with TFE as the proton source. CPE experiments in the presence of TEAHPF₆ showed a CO efficiency of $83 \pm 1\%$ with a TOF of 0.87 s⁻¹ for complex 1, whereas complex 2 demonstrates quantitative FE_{CO} (104 ± 4%) with TEAHPF₆, and an increased TOF of 13.70 s⁻¹. Finally, a CPE experiment was conducted with 0.1 M TEAHPF₆ serving the dual role of electrolyte and proton source for 2 demonstrated quantitative FE_{CO} and a TOF of 5.47 s⁻¹. When TEAHPF6 is used as electrolyte and proton source, more positive potentials are required to suppress background HER, suggesting that the TBA⁺ component of the electrolyte coulombically screens the strength of the interaction between the electrode and TEAH⁺, improving the molecular electrocatalytic response.

Computational results

Given the observed activity difference for complexes 1 and 2, several of the key reaction steps were evaluated to see if this correlated to an intrinsic difference, Table 2. Using a DFT methodology developed in previous comprehensive computational studies,23 a minimal difference is predicted for the thermodynamics and kinetics of CO2 binding, the thermodynamics and kinetics for the rate-limiting protonation of the [Cr-CO₂H]⁻ intermediates by TEAH⁺, as well as for the thermodynamics of CO release. Calculation of the redox potential for the generation of the catalytically active species for complex 1, $\left[\operatorname{Cr}(^{p-tbu}dhbpy)\right]^{0/-} = -1.96 \text{ V } \nu s. \text{ Fc}^+/\operatorname{Fc}$, showed good agreement with experimental value of -1.95 V. Likewise, the calculated redox potential at which the catalytically relevant species is generated for complex 2, $[Cr(^{nPr}dhbpy)]^{0/-} = -$ 1.97 V vs. Fc⁺/Fc, was similarly aligned with the experimental value of -1.95 V.

Discussion

Using the mechanistic data described above, it is possible for us to propose a role for the pendent relay in modifying the intrinsic catalytic mechanism (Fig. 1 and eqn (1)-(6)). It is helpful here to first summarize the catalytic cycle of the base catalyst family as the following equilibrium and irreversible reaction steps (note for simplicity a neutral acid, HA, is considered, solvento species are not identified, and that the ligand, which remains tetradentate in all cases, is omitted):

$$[Cr]^{0} + e^{-} \rightleftharpoons [Cr]^{-} \quad E_{cat}^{\circ} \tag{1}$$

$$[Cr]^- + CO_2 \; \rightleftharpoons \; [Cr(CO_2)]^- \; \textit{K}_{CO_2} \tag{2}$$

$$\left[Cr(CO_2)\right]^- + HA \rightarrow \left[Cr(CO_2H)\right]^0 + A^- \quad \textit{k}_1 \eqno(3)$$

$$[Cr(CO_2H)]^0 + e^- \rightleftharpoons [Cr(CO_2H)]^- \quad E_1^{\circ} \tag{4}$$

$$[Cr(CO_2H)]^- + HA \rightarrow [Cr(CO)]^0 + H_2O + A^- \quad k_2$$
 (5)

$$[Cr(CO)]^0 \rightleftharpoons [Cr]^0 + CO K_{CO}$$
 (6)

Previous mechanistic and computational studies have established that E_1° is more positive than E_{cat}° , that K_{CO_2} is an endergonic process (<1), that CO release is facile (no CO adducts observed, $K_{CO} \gg 1$), and that $k_1 \gg k_2$. Overall, this allows the electrocatalytic behavior of the original catalyst to be condensed to an ECEC' framework, considering the two proton transfers to be the relevant chemical steps. Consistent behavior is observed here for complexes 1 and 2, where no CO₂ binding is again observed under aprotic conditions (Fig. 3). However, several notable changes occur in conditions with CO2 and a proton donor present. In many cases, mixed-order kinetics are observed for the substrates and the metal complex. Further, in the case of complex 2, a shift to more positive voltages is observed for the catalytic potential with two of the three proton donors. Since the catalytically relevant reduction potentials of 1 and 2 are nearly identical, it is proposed that several pre-equilibrium reactions are contributing to the deviation of both 1 and 2 from the previously reported catalytic behavior (eqn (7) and (11)), as summarized by the modified reaction sequence below:

$$[Cr]^0 + e^- \rightleftharpoons [Cr]^- \quad E_{cat}^{\circ} \tag{1}$$

$$[Cr]^- + HA \rightleftharpoons Cr(HA)]^- K_{HA1}$$
 (7)

$$[Cr(HA)]^- + CO_2 \rightleftharpoons [Cr(CO_2)(HA)]^- K_{HA_CO_2}$$
 (8)

$$[Cr(CO_2)(HA)]^- \rightarrow [Cr(CO_2H)]^0 + A^- k_1$$
 (9)

$$\left[\operatorname{Cr}(\operatorname{CO}_{2}\operatorname{H})\right]^{0} + e^{-} \rightleftharpoons \left[\operatorname{Cr}(\operatorname{CO}_{2}\operatorname{H})\right]^{-} \quad E_{1}^{\circ} \tag{10}$$

$$[Cr(CO_2H)]^- + HA \rightleftharpoons [Cr(CO_2H)(HA)]^- K_{HA2}$$
 (11)

$$[Cr(CO_2H)(HA)]^- \rightarrow [Cr(CO)]^0 + H_2O + A^- \quad k_2$$
 (12)

Table 2 Summary of DFT results for key reaction steps

Reaction	ΔG^{\ddagger} (kcal mol ⁻¹)	ΔG (kcal mol ⁻¹)
$[Cr(^{p-tbu}dhbpy)]^{-} + CO_{2} \rightleftharpoons [Cr(^{p-tbu}dhbpy)(CO_{2})]^{-}$	+11.1	+6.6
$[Cr(^{nPr}dhbpy)]^- + CO_2 \rightleftharpoons [Cr(^{nPr}dhbpy)(CO_2)]^-$	+10.9	+6.5
$[Cr(P^{-tbu}dhbpy)(CO_2H)]^- + TEAH^+ \rightarrow [Cr(P^{-tbu}dhbpy)(CO)]^0 + H_2O + TEA$	+10.5	-19.4
$[Cr(^{nPr}dhbpy)(CO_2H)]^- + TEAH^+ \rightarrow [Cr(^{nPr}dhbpy)(CO)]^0 + H_2O + TEA$	+11.9	-19.0
$[Cr(P^{-tbu}dhbpy)(CO)]^0 \rightleftharpoons [Cr(P^{-tbu}dhbpy)]^0 + CO$		-7.4
$[Cr(^{nPr}dhbpy)(CO)]^0 \Rightarrow [Cr(^{nPr}dhbpy)]^0 + CO$		-7.8

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$$[Cr(CO)]^0 \approx [Cr]^0 + CO K_{CO}$$
 (6)

The data suggest that for both proton transfer steps (k_1 and k_2), a pre-equilibrium involving the association of the acid to reduced form of complex 2, $K_{\rm HA1} > 1$, becomes kinetically relevant in the presence of certain acids. It is possible to extract an equilibrium binding constant for the equilibrium interaction between 2 and TEAH $^+$ under Ar saturation of $K_{\rm HA1} = 380 \pm 50$ (Fig. S64 $^+$) at the reduction feature where catalysis is initiated under CO $_2$ saturation. A comparable shift in reduction potential under Ar saturation is not observed for 2 with the weaker and uncharged PhOH and TFEOH proton donors or complex 1 with any of the three proton donors.

The shift observed for the $E_{\rm cat/2}$ of complex 2 suggests that these pre-equilibria are catalytically relevant when the methoxy group is present. In a reductive ECEC' framework, where the second reversible electron transfer occurs at potentials more positive than the first, shifts in catalytic potential can be described according to the following equation: 66

$$E_{1/2} = E_{\rm P/Q}^{\circ} + \frac{RT}{F} \ln \left(1 + \frac{\sqrt{k_1 C_{\rm A}^{\circ}}}{\sqrt{k_2 C_{\rm Z}^{\circ}}} \right)$$
 (13)

In this framework, positive shifts in potential will be observed as the second-order rate constant k_1 increases relative to k_2 . For the complexes described here, C_A° is taken as the concentration of the proton donor and the TOF derived from preparative electrolysis is assigned as $k_2C_2^{\circ}$, given that k_2 is the rate-limiting step of the reaction and that CVs of the catalytic waves under these conditions have the appropriate S-shape for a kinetically limited response and experience minimal change in current as proton donor concentration is varied. Using the relationship of $E_{\text{cat/2}}$ and acid concentration, a k_1 of 11 ± 2 M⁻¹ s⁻¹ was determined with TFE as a proton donor by averaging values across a concentration range of 0.1 to 1.1 M TFE (Fig. S63†).66 However, comparable analysis of the shifts in $E_{\text{cat/2}}$ of 2 in relation to acid concentration under electrocatalytic conditions showed that the same k_1 value steadily increased from $7.1 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$ at 20 mM TEAH to 2.7×10^4 M⁻¹ s⁻¹ at a final concentration of 150 mM (Fig. S65†).⁶⁷ It is reasoned that the cationic charge introduced in the secondary coordination sphere upon the association of TEAH⁺ to the anionic Cr complex contributes favorable kinetic and thermodynamic effects to the conversion of reduced CO₂ species in a manner analogous to that previously observed for other complexes. 15,16,68-74 It is important to acknowledge, however, that it is difficult to completely disentangle this effect from the increased proton donor activity of TEAH⁺ relative to TFE. As mentioned above, there was no shift in $E_{\text{cat/2}}$ observed for PhOH with complex 2 and no shift in $E_{cat/2}$ for complex 1 with any of the three proton donors. Consistent with the proposed relevance of pre-equilibrium steps when TEAH⁺ is used as the acid, there is an absence of any significant changes in the energetics and barriers of the key reaction steps for complexes 1 and 2 by DFT, Table 2.

A pre-equilibrium condition can also be used to rationalize the observed increase in catalytic activity observed for complex 2 relative to complex 1 in preparative electrolysis, as outlined recently by Pattanayak and Berben.⁵⁹ The experimental observations can also be contrasted with the Fe porphyrin derivative reported by Savéant and co-workers, who showed that K_{CO2} was impacted by the inclusion of pendent proton donors, thanks to intramolecular hydrogen-bonding interactions. 47,56 For the Fe porphyrin derivative with pendent proton donors reported by Savéant and co-workers, hydrogen bonding stabilized the Fe-CO2 intermediate, shifting the corresponding redox feature to more positive potentials, however, the subsequent reduction that initiates the rate-determining proton-coupled electron transfer was not shifted by a similar amount, resulting in the appearance of a pre-wave to the catalytic feature. It is reasoned that the difference in the impact of the pendent relay site on reaction parameters reported here has to do with the position of the pendent relay relative to the site of CO₂ binding.

A plot of $log_{10}(TOF s^{-1})$ versus $E_{cat/2}$ for the original family of Cr-based molecular catalysts, 10,19,21 with PhOH, for CO2 to CO shows a normal Tafel relationship (black squares, Fig. 5), where more negative potentials show an increased TOF. Comparing this trend with the data collected for 1 and 2 with PhOH (blue and red square, respectively, Fig. 5), the more positive $E_{\text{cat/2}}$ of both species corresponds to a decreased TOF, consistent with a Tafel scaling relationship (more negative catalytic potentials result in greater activity for the same catalyst family).⁷⁵ However, the response of complex 1 in this series is significantly below the linear trend established by the original complexes and 2. This deviation suggests that the propensity to reversible electrode adsorption limits the activity of 1. Although estimated to be slightly a weaker acid, 63,76 TFE shows an increase in activity for both 1 and 2 relative to PhOH, with a slight positive shift in $E_{\text{cat/2}}$ observed for the latter. These data lead to the assumption that there are additional contributions

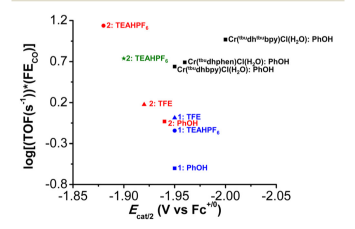


Fig. 5 Comparison of activity and standard reduction potential in the Cr catalyst family. Squares are data taken with PhOH and added supporting electrolyte. Triangles are data taken with TFE and added supporting electrolyte. Circles are data taken with triethylammonium as the acid and added supporting electrolyte, while the star indicates data taken with triethylammonium in the absence of supporting electrolyte.

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to the effective proton activity of TFE, or that there are limitations to the estimated pK_a . It is possible that the difference may have to do with a combination of solvation effects, homoconjugation, and binding of CO₂ by the conjugate base of TFE, since the pK_a of TFE is higher than PhOH in known solvents. 63,77

In the presence of the stronger acid, TEAHPF₆ (blue and red circles), a significant separation in $E_{\text{cat/2}}$ is observed between 1 and 2 (60 mV), with the more positive potential of complex 2 showing significantly higher activity. Since an equilibrium binding constant can be established for complex 2 but not 1 under Ar saturation with this acid (Fig. S64†), this effect must be correlated to the involvement of the methoxy group in proton transfer steps following CO₂ binding at Cr. Further, the sensitivity of $E_{\text{cat/2}}$ to the concentration of the ammoniumbased acid can be used to demonstrate that the second-order rate constant of the underlying proton transfer in question, k_1 , has increased significantly from TFE (Fig. S63 and S65†). Thus, it appears that the proton donor can render the pre-equilibrium steps to the formation of intermediate iii kinetically relevant with suitable functional groups in the secondary coordination sphere (Fig. 1).

The potential tuning effect resulting from modifying this pre-equilibrium between complexes 1 and 2 is also noteworthy, especially in light of their activity differences. While TEAH⁺ associates to the anionic complex i derived from 2, hydrogenbonding interactions with the pendent methoxy groups would not stabilize CO₂ binding based on relative positioning. However, for coulombic reasons, association of the ammonium proton donor to anionic states is overall favorable and observed to improve proton transfer to the electronegative O atoms of bound CO2 reduction intermediates, as negative charge develops there during C=O bond breaking. Indeed, Martin and Mayer demonstrated that it is the amount of positive charge, rather than the position of the charge, which matters in activity enhancements.16 However, the utility of using cationic acids to achieve this is tempered by the significant increase in overpotential required. Going forward, these data imply that in order to harness this improved operating potential, the pre-equilibrium proton donor association needs to be enhanced when weaker acids are used. This suggests that optimizing the pendent relay ligand design, rather than proton donor choice, will ultimately lead to the most significant performance increases in the long-term.

While pendent functional groups increasing activity at a fixed reduction potential are known48,78 and the pre-equilibrium effect has been previously observed to impact speciation during electrocatalysis,59 future efforts could focus on the combination of these effects to achieve inverse potential scaling for homogeneous electrocatalytic activity.⁷⁹ It is worth noting that alternative strategies for accessing inverse potential scaling based on the ligand and reaction system design also exist and offer complementary opportunities to tune performance. 10,80 Since there has been general interest in the introduction of charge to impact the kinetics and operating potential of molecular CO2 reduction catalysts, 15,16,80 in

addition to the interest in pendent relays described above, it would be advantageous to leverage both effects.⁷⁹

Conclusion

These experimental results demonstrate that the combination of a pre-equilibrium effect directed by a pendent functional group and the use of a cationic proton donor can achieve improved activity at lower operating potentials during homogeneous CO2 reduction. Mechanistic studies comparing neutral proton donors to the cationic one show minimal changes in activity or operating potentials unless the cationic acid is present, with a particularly strong interaction occurring in the presence of the pendent relay. Therefore, it is reasoned that the use of cationic acids can generally be regarded as beneficial when the catalytic reduction cycle involves several anionic intermediate species in organic solvents. Based on these conclusions, further studies exploring cationic acids and optimizing the ligand framework to interact with the metal complex following CO2 binding and activation are currently underway. Importantly, this strategy should be applicable to other proton-coupled electron transfer steps in small molecule reduction reactions, offering a new design strategy for optimizing activity and selectivity.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no competing financial interest.

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