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# Directing the reactivity of metal hydrides for selective CO<sub>2</sub> reduction

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A critical challenge in electrocatalytic CO<sub>2</sub> reduction to renewable fuels is product selectivity. Desirable products of CO2 reduction require proton equivalents, but key catalytic intermediates can also be competent for direct proton reduction to H<sub>2</sub>. Understanding how to manage divergent reaction pathways at these shared intermediates is essential to achieving high selectivity. Both proton reduction to hydrogen and CO<sub>2</sub> reduction to formate generally proceed through a metal hydride intermediate. We apply thermodynamic relationships that describe the reactivity of metal hydrides with  $\mathrm{H}^+$  and  $\mathrm{CO}_2$  to generate a thermodynamic product diagram, which outlines the free energy of product formation as a function of proton activity and hydricity ( $\Delta G_{H-}$ ), or hydride donor strength. The diagram outlines a region of metal hydricity and proton activity in which CO<sub>2</sub> reduction is favorable and H<sup>+</sup> reduction is suppressed. We apply our diagram to inform our selection of [Pt(dmpe)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> as a potential catalyst, because the corresponding hydride [HPt(dmpe)<sub>2</sub>]<sup>+</sup> has the correct hydricity to access the region where selective CO<sub>2</sub> reduction is possible. We validate our choice experimentally; [Pt(dmpe)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> is a highly selective electrocatalyst for CO<sub>2</sub> reduction to formate (>90% Faradaic efficiency) at an overpotential of less than 100 mV in acetonitrile with no evidence of catalyst degradation after electrolysis. Our report of a selective catalyst for CO<sub>2</sub> reduction illustrates how our thermodynamic diagrams can guide selective and efficient catalyst discovery.

electrocatalysis | CO2 reduction | solar fuel | formate production | hydride

The emerging availability of inexpensive renewable electricity has motivated interest in using electrolytic methods to generate sustainable fuels. Electrocatalytic  $CO_2$  reduction provides an entry to carbon-neutral fuels, but product selectivity remains a significant challenge (1, 2). Nearly all reductive reactions of interest involve protons as well as electrons, introducing the complication of direct proton reduction to H<sub>2</sub> under electrolytic conditions. Diversion of electron equivalents into proton reduction results in lower Faradaic efficiency for the desired  $CO_2$ reduction reaction. Various strategies to inhibit or suppress H<sub>2</sub> evolution for heterogeneous (3–7) and homogeneous (8, 9) catalysts have been explored.

To understand the factors that determine selectivity between  $CO_2$  and H<sup>+</sup> reduction, we have been investigating the reactivity of metal hydrides. Selectivity for formate production is particularly challenging because metal hydride intermediates are common to both reaction pathways (Fig. 1). As a result, very few heterogeneous (10–13) or homogeneous (14–16) catalysts have been reported with high (>90%) Faradaic efficiency for formate production. Understanding the reactivity of metal hydrides is key to controlling the bifurcating reaction pathways that ultimately determine selectivity. Most selective catalysts for  $CO_2$  reduction utilize kinetic inhibition (or a high-transition state barrier) to minimize H<sub>2</sub> evolution.

In this report, we utilize a thermodynamic approach to describe the reactivity of metal hydrides with H<sup>+</sup> and CO<sub>2</sub> by applying known free energy relationships (14, 17–28). We depict our findings in a diagram (Fig. 2) that describes the thermodynamic products as a function of hydricity ( $\Delta G_{H-}$ ) (Eq. 1) of the metal

hydride intermediate vs. proton activity ( $pK_a$ ). Notably, a region is delineated in which select combinations of hydricity and  $pK_a$  lead to thermodynamic favorability for selective CO<sub>2</sub> reduction to formate over hydrogen evolution. Since hydricity values have been measured for many classes of compounds and can be predictably tuned through metal and ligand design (29), we believe that the diagrams provide instructive guidelines for targeting the discovery of selective reduction catalysts.

We establish the utility of our diagram by using it to select a known metal hydride  $[HPt(dmpe)_2]^+$  with an appropriate hydricity in acetonitrile to access conditions in which selective CO<sub>2</sub> reduction to formate is favorable. We demonstrate that the resting state catalyst  $[Pt(dmpe)_2](PF_6)_2$  reduces CO<sub>2</sub> to formate with high Faradaic efficiency and low overpotential.

### **Results and Discussion**

**Thermodynamic Product Diagrams.** We first describe the construction of the thermodynamic product diagram in acetonitrile (CH<sub>3</sub>CN), but the relationships hold in other solvents provided that the correct constants are used (*vide infra*). The thermodynamic cycle describing the free energy for the reaction of a metal hydride toward H<sup>+</sup> to evolve H<sub>2</sub> is shown in Fig. 2. The free energy of H<sub>2</sub> evolution (Eq. 4) is dependent on the hydricity, pK<sub>a</sub> of the acid, and the heterolytic bond forming energy of H<sub>2</sub>, which is 77 kcal/mol in acetonitrile (30). Eq. 4 has successfully been applied to identify and optimize electrocatalysts for H<sub>2</sub> evolution (26, 31– 38) and H<sub>2</sub> oxidation (31–34, 39–42), and to achieve reversible reactivity (43, 44).

The relationship between  $pK_a$ , hydricity, and  $H_2$  evolution in acetonitrile is quantitatively depicted in Fig. 24. The black line

### Significance

Electrocatalytic CO<sub>2</sub> reduction to fuels using renewable electricity provides an entry into sustainable carbon-neutral fuels. Most reactions require proton equivalents, but direct proton reduction to H<sub>2</sub> leads to low product selectivity, a significant challenge in the field. The issue of product selectivity arises from shared catalytic intermediates for both CO<sub>2</sub> reduction and H<sub>2</sub> evolution. We describe how thermodynamic relationships regarding the reactivity of H<sup>+</sup> and CO<sub>2</sub> with one of these shared intermediates, homogeneous metal hydrides, can be applied to identify catalyst candidates and conditions where H<sub>2</sub> evolution is suppressed in favor of highly selective CO<sub>2</sub> reduction. The conceptual framework is used to outline operational conditions for electrocatalytic reduction of CO<sub>2</sub> to formate (HCO<sub>2</sub><sup>-</sup>) by [Pt(dmpe)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> with negligible H<sub>2</sub> production.

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**Fig. 1.** Proposed catalytic cycles for H<sup>+</sup> and CO<sub>2</sub> reduction.

differentiates the boundary in which the free energy of Eq. 4 in Fig. 2 ( $\Delta G_{H2}$ ) equals zero and MH/H<sup>+</sup>/H<sub>2</sub> exist in equilibrium, analogous to lines in a Pourbaix diagram. Protonation to evolve hydrogen is exergonic for hydride complexes of a given hydricity under conditions below the black line (green zone) and endergonic above the line (blue zone).

The thermodynamic requirement for reduction of CO<sub>2</sub> to formate, a net hydride transfer, is also dictated by the hydricity ( $\Delta G_{H-}$ ) of the donor as shown in Eq. **6** (Fig. 2). Eq. **6** has been applied to rationalize or predict the activity of the iron-based CO<sub>2</sub> reduction electrocatalyst by Berben and coworkers (20) and CO<sub>2</sub> hydrogenation catalysts (45–50) as well as formate oxidation electrocatalysts (51). Transition metal hydricity values lower than that of formate will result in exergonic hydride transfer to CO<sub>2</sub>. This information can be mapped onto the thermodynamic product diagram illustrating metal hydride reactivity with protons (Fig. 2*B*). The thermodynamic product distribution that results from stoichiometric mixtures of metal hydride and acid of specific  $pK_a$  values under 1 atm of CO<sub>2</sub> is described in this diagram. The  $pK_a$  of formic acid is estimated to be ~20.9 in acetonitrile (52); therefore, it is expected to be protonated at lower  $pK_a$  values.

In Fig. 2B, a region is defined where specific metal hydricity and  $pK_a$  combinations will result in selective reduction of CO<sub>2</sub> to formate without concomitant H<sub>2</sub> evolution (near the bottom right of Fig 2B). The challenge of selective  $CO_2$  reduction is often introduced by comparing the thermodynamic potentials for  $H^+$  reduction with CO<sub>2</sub> reduction to formate [-0.028 V (53) vs. -0.150 V (22) vs.  $Fe(C_5H_5)_2^{+/0}$  in CH<sub>3</sub>CN]. Since the former is more positive than the latter, hydrogen is assumed to be the more favorable thermodynamic product upon reduction. However, these potentials are listed at the standard state (1 M H<sup>+</sup>). A full Pourbaix diagram that spans a larger range of proton activity  $(pK_a)$  provides a more complete picture (SI Appendix, Fig. S1). The thermodynamic potential for proton reduction shifts with the  $pK_a$  of the solution by 59 mV per unit according to the Nernst equation for a 2 e<sup>-</sup>, 2H<sup>+</sup> process. Similarly, CO<sub>2</sub> reduction to formic acid follows the same relationship until the solution  $pK_a$  matches that of formic acid to generate formate. For CO<sub>2</sub> reduction to formate, the thermodynamic potential shifts will deviate in accordance with a 2 e<sup>-</sup>, 1H<sup>+</sup> process or 29.5 (59/2) mV per  $pK_a$  unit. The smaller decline in thermodynamic potential vs.  $pK_a$  compared with H<sup>+</sup> reduction results in a crossing of potentials, where  $CO_2$  reduction to formate occurs at a more positive potential at higher  $pK_a$  values and can thus be

the more favorable thermodynamic product. The crossing point in the Pourbaix diagram (*SI Appendix*, Fig. S1) matches the  $pK_a$ (25.1), which defines the initial point in which selective CO<sub>2</sub> reduction can be achieved in our diagram. A similar crossing in the 2H<sup>+</sup>/H<sub>2</sub> and CO<sub>2</sub>/HCO<sub>2</sub><sup>-</sup> is observed in Pourbaix diagrams in water (54), although it is complicated by CO<sub>2</sub> equilibria with hydroxide at high pH (55). We believe that our redrawn thermodynamic product diagram, where potential is replaced by hydricity, provides a more instructive guide for targeting catalysts







**Fig. 2.** Free energy calculations for  $H_2$  and  $HCO_2^-$  generation from metal hydrides and aqueous thermodynamic product diagrams (A and B).

to access the region where selective  $CO_2$  reduction is possible, which we demonstrate experimentally herein.

A recently published perspective by Kubiak and coworkers (22) described a primarily linear relationship between hydricity and the one-electron reduction potentials of the parent complex for previously reported groups 8-10 metal hydrides. The perspective provides valuable insight into the relationship between hydricity and reversible redox properties as they pertain to  $H_2$ evolution and CO<sub>2</sub> reduction. However, the authors note that their correlation is qualitative and dependent on similar hydrogen atom transfer bond dissociation free energies among their complexes. Additionally, the authors detail why some metal hydrides deviate significantly from their observed trend and were excluded from the linear fit. We note that the diagrams shown in Fig. 2 are based directly on thermodynamic hydricity values and are thus applicable to all metal hydrides, regardless of metal identity, oxidation state, or ligand environment. The thermodynamic hydricity can also be accurately measured for complexes without reversible redox potentials using methods that have previously been described (19).

Application in other solvents. The diagrams shown in Fig. 2 can be drawn for any solvent provided that the appropriate constants  $(\Delta G_{H2} \text{ and } \Delta G_{HCO2-})$  are known in the respective solvent. For example, the diagram illustrating the relationships between metal hydricity and proton activity (similar to Fig. 24) in dimethylsulfoxide (DMSO) is shown in *SI Appendix*, Fig. S2 ( $\Delta G_{H2} = 60.7$  kcal/mol in DMSO) (30). The hydricity of formate ( $\Delta G_{HCO2-}$ ) in DMSO is estimated to be 42 kcal/mol (56). The equivalent diagram shown in Fig. 2B for DMSO is shown in SI Appendix, Fig. S3. We note that we were unable to find a published value for the  $pK_a$  of formic acid in DMSO, and therefore, the line between  $CO_2/HCO_2^{-1}$  is drawn across the whole range of pK<sub>a</sub> values shown. Applying thermodynamic relationships with metal hydricity is most useful in organic solvents with self-consistent data on acid  $pK_a$  values. Most measurements of this type have been made in CH<sub>3</sub>CN (57–61) and to a lesser extent DMSO (62), but additional values in various solvents are continually being reported.

 $\Delta G_{H2}$  and  $\Delta G_{HCO2-}$  have also been reported in aqueous solution (34.2 and 24.1 kcal/mol, respectively) (17). The analogous diagrams in Fig. 2*A* and *B* are shown in *SI Appendix*, Figs. S4 and S5, respectively. However, the diagrams do not account for the equilibria between CO<sub>2</sub> and OH<sup>-</sup> to form HCO<sub>3</sub><sup>-</sup> and CO<sub>3</sub><sup>2-</sup>, which become more significant at higher pH values.

The hydricity value ( $\Delta G_{H-}$ ) for metal hydrides also has a solvent dependence. Multiple studies have measured how solvent affects the hydricity values (14, 18, 19, 23–25, 27, 35). Hydricity values for transition metal hydrides tend to be lower and span a narrower range in solvents with higher dielectric constants, but their absolute values do not change in a predictable fashion by solvent. However, a notable trend is that hydricity values for transition metal hydrides in organic solvents tend to decline to a greater magnitude (become better donors) in water compared to  $HCO_2^{-1}$  ( $\Delta G_{HCO2-}$ ). As a result, there are a few examples where  $CO_2$  reduction to  $HCO_2^{-1}$  is exergonic in water but endergonic in organic solvents (14, 27, 48). Additionally, detailed studies by Miller and coworkers (63) determined that aqueous hydricity can also be dependent on anions commonly found in aqueous buffers as well as hydroxide at higher pH values.

*Kinetic vs. thermodynamic selectivity.* The diagrams depicted in this study delineate how to achieve thermodynamic selectivity for CO<sub>2</sub> reduction. In CH<sub>3</sub>CN, DMSO, and H<sub>2</sub>O, thermodynamic selectivity can only be achieved at modest or low proton activity ( $pK_a > 25.9$  in CH<sub>3</sub>CN,  $pK_a > 13.6$  in DMSO, and pH > 8.1 in H<sub>2</sub>O). However, kinetic barriers for protonation can be applied to achieve selectivity for CO<sub>2</sub> reduction under more acidic conditions. The difference in kinetic selectivity will still have to compensate for larger free energies of protonation of the metal hydride to form hydrogen at lower  $pK_a/pH$  values. Consistent with

the free energy associated with  $H_2$  and  $HCO_2^-$  formation (Eqs. 4 and 7, respectively) the only two molecular catalysts with >90% selectivity for formate generation function optimally at modest to high  $pK_a/pH$  conditions, with decreasing selectivity at lower proton activities (16, 20). Yet, both maintain fairly high selectivity under more acidic conditions, indicating that the product distribution is also under kinetic control (64).

Application of Thermodynamic Product Diagrams for Catalyst Discovery. To target the region in Fig. 2*B* where selective  $CO_2$  reduction to  $HCO_2^-$  in  $CH_3CN$  is dictated by the thermodynamic Eqs. 4 and 6, we selected  $[Pt(dmpe)_2](PF_6)_2$  (3) as a possible catalyst. The corresponding hydride  $[HPt(dmpe)_2](PF_6)$  (2) has an experimentally measured hydricity ( $\Delta G_{H-}$ ) (Eq. 1) of 41.4 kcal/mol (Fig. 2) (19, 65). Consistent with Eq. 6,  $[HPt(dmpe)_2](PF_6)$  reacts stoichiometrically with  $CO_2$  to give  $HCO_2^-$  and  $[Pt(dmpe)_2](PF_6)_2$  (3) (21). The preparation and characterization of the hydride precursor  $Pt(dmpe)_2$  (1) along with 2 and 3 have been previously reported (65, 66). Complexes 1, 2, and 3 are diamagnetic with easily distinguishable  ${}^{31}P{}^{1}H$  NMR spectral signatures.

According to Eq. 4 and Fig. 2, the threshold for H<sub>2</sub> evolution with [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (2) is at a pK<sub>a</sub> of 25. Acids with pK<sub>a</sub> values lower than 25 will be exergonic for H<sub>2</sub> evolution. As expected, stoichiometric reaction of [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (2) with acids, such as anilinium tetrafluoroborate (pK<sub>a</sub> = 10.62) and protonated 1,8-diazabicyclo[5.4.0]undec-7-ene (pK<sub>a</sub> = 24.34) in CH<sub>3</sub>CN (60), results in protonation to give H<sub>2</sub> and [Pt(dmpe)<sub>2</sub>] (PF<sub>6</sub>)<sub>2</sub> (3) (*SI Appendix*, Figs. S6 and S7). In contrast, [HPt (dmpe)<sub>2</sub>](PF<sub>6</sub>) (2) is stable to protonation using acids with higher pK<sub>a</sub> values ( $\Delta G_{H2} > 0$ ), which we demonstrate experimentally. No reaction is observed with [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (2) upon addition of phenol (pK<sub>a</sub> = 29.1) (*SI Appendix*, Fig. S8) (62). According to Eq. 4, H<sub>2</sub> evolution with [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (2) and phenol has an endergonic free energy of 5.5 kcal/mol.

The  $pK_a$  (31.1) (65) of the metal hydride provides an upper bound on acids that can be used for metal hydride generation. As a result, [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (**2**) can be selectively generated at an electrode using proton sources with  $pK_a$  values between 31.1 and 25. Therefore, phenol ( $pK_a = 29.1$ ) is sufficiently acidic to generate [HPt(dmpe)<sub>2</sub>](PF<sub>6</sub>) (**2**) without additional protonation to evolve H<sub>2</sub>.

Selective electrocatalytic reduction of  $CO_2$  to formate. [Pt(dmpe)\_2] (PF<sub>6</sub>)<sub>2</sub> (3) exhibits a 2 e<sup>-</sup> reduction to [Pt(dmpe)\_2] (1) at -1.73 V vs. Fe(C<sub>5</sub>H<sub>5</sub>)<sub>2</sub><sup>+/0</sup> in acetonitrile (Fig. 3, black trace). Electrochemical reduction of 3 to 1 in the presence of phenol results in a loss of reversibility (blue trace in Fig. 3), which we attribute to formation of the hydride 2 [electron transfer (E) followed by a chemical step (C), protonation, or an EC step]. Higher concentrations of phenol do not result in an increase in current (green trace in Fig. 3).

According to Eq. 5,  $[HPt(dmpe)_2](PF_6)$  (2) is sufficiently hydritic to react with CO<sub>2</sub> to generate formate ( $\Delta G^\circ$  in Eq. 6 = -2.6 kcal/mol). The cyclic voltammogram on addition of 1 atm of CO<sub>2</sub> to 3 with phenol is shown by the red trace in Fig. 3. Titration with increasing concentrations of phenol results in an increase in current (Fig. 3), which is indicative of electrocatalysis. The current reaches a maximum at 10 equivalents.

A proposed catalytic cycle for CO<sub>2</sub> reduction to formate with thermodynamic parameters ( $E_{1/2}$  for e<sup>-</sup> transfer and pK<sub>a</sub> for protonation reactions) for each step is depicted in Fig. 4. Electrolysis was performed at -2.4 V vs. Fe(C<sub>5</sub>H<sub>5</sub>)<sub>2</sub><sup>+/0</sup> under 1

Electrolysis was performed at -2.4 V vs. Fe(C<sub>5</sub>H<sub>5</sub>)<sub>2</sub><sup> $\pm$ /0</sup> under 1 atm of CO<sub>2</sub> (1 mM **3** in 10 mM phenol) for 1 h (*SI Appendix*, Figs. S9 and S10). The concentration of formate in solution after electrolysis was measured using two different methods. The first method used an internal dimethylforamide (DMF) standard to quantify formate by <sup>1</sup>H NMR spectroscopy (*SI Appendix*, Fig. S11). The second method required acidifying the postelectrolysis



**Fig. 3.** Cyclic voltammetry under 1 atm of N<sub>2</sub> of 0.5 mM solution of [Pt (dmpe)<sub>2</sub>](PF<sub>6</sub>)<sub>2</sub> (**3**) (black); after addition of 1 equivalent of phenol (blue); and 5 equivalents of phenol (green); and under 1 atm of CO<sub>2</sub> with 5 equivalents of phenol (red); and 10 equivalents of phenol (orange). Conditions: 0.1 M Et<sub>4</sub>NPF<sub>6</sub>, 1 mM Fe(C<sub>5</sub>H<sub>5</sub>)<sub>2</sub> present as an internal reference, glassy carbon working and auxiliary electrode, Ag/AgCl pseudoreference electrode, 10 mV/s scan rate.

solution with HCl and quantifying the formic acid with a DMF internal standard using <sup>1</sup>H NMR spectroscopy (*SI Appendix*, Fig. S12). The Faradaic efficiency for formate production from either method was greater than 90%. The headspace of the cell was analyzed by GC to detect and quantify H<sub>2</sub> and CO production. A small amount of H<sub>2</sub> (<0.1%) was sometimes observed, but CO was never detected. <sup>31</sup>P{<sup>1</sup>H} NMR spectra of the pre- and postelectrolysis solution display a single resonance that corresponds to **3**. Quantification by integration relative to the PF<sub>6</sub><sup>-</sup> anion (*SI Appendix*, Figs. S13 and S14) confirms retention of the catalyst after electrolysis.

The maximum catalytic current observed in the cyclic voltammetry (Fig. 3, orange trace) and the peak current under noncatalytic conditions (Fig. 3, black trace) was used to estimate an observed rate constant ( $k_{obs}$ ) of 0.5 s<sup>-1</sup> (*SI Appendix*, Eq. S4) (67, 68) for CO<sub>2</sub> reduction to formate. The standard potential for CO<sub>2</sub> reduction to formate in acetonitrile was recently estimated (22). The thermodynamic potential under our conditions ( $pK_a$  of 29.1) is E° = -1.64 V. Using the half-wave potential E°<sub>1/2</sub> = -1.73 V (69), the overpotential for catalysis is 90 mV.

To quantify any potential  $H_2$  or formate generated directly at the electrode, an equivalent electrolysis in the absence of any Pt compound was performed (*SI Appendix*, Fig. S15). Analysis of the headspace by GC detected the presence of <0.1% mL  $H_2$ , no CO, and no formate in solution by <sup>1</sup>H NMR spectroscopy. Calibration curves for  $H_2$  and formate quantification are shown in *SI Appendix*, Figs. S16 and S17, respectively.

**General considerations.** The diagrams in Fig. 2 informed our choice of  $[Pt(dmpe)_2](PF_6)_2$  (3) as a good catalyst candidate for selective CO<sub>2</sub> reduction to formate. The hydricity of the corresponding hydride  $[HPt(dmpe)_2](PF_6)$  (2) is sufficient to reduce CO<sub>2</sub>, and there are appropriate acids to access the region of selective reduction. Although the thermodynamic framework described herein proved effective for  $[Pt(dmpe)_2]^{2+}$  (3), some considerations on its application to other potential catalysts are discussed.

Although nearly all molecular hydrogen evolution reaction (HER) catalysts involve a metal hydride intermediate, H–H bond formation can occur through two distinct mechanisms: bi-molecular (homolytic) or heterolytic (protonation; as described

here). There are cases where both homolytic and heterolytic H<sub>2</sub>forming pathways are viable, such as in the well-studied cobaloxime HER catalysts (70–72), with the predominant operating mechanism dependent on the specific reaction conditions (73–80). The free energy of homolytic H–H bond formation is dependent on the homolytic bond energy of the metal hydride and H<sub>2</sub> and is not expected to have a dependence on the proton activity.

Consequently, the thermodynamic framework described in Fig. 2 is most useful for metal hydrides that are stable to homolytic H<sub>2</sub> bond formation. This is generally true for many classes of transition metal hydrides, including those in the  $[Ni(P_2N_2)_2]^{2+}$  class of catalysts (70, 71, 78–91), the water soluble  $[HNi(DHMPE)_2]^{2+}$ , and the subject of these studies  $[HPt(dmpe)_2](PF_6)_2$  (2). [The <sup>31</sup>P{<sup>1</sup>H}NMR spectra of  $[HPt(dmpe)_2](PF_6)_2$  (2) in acetonitrile show no change over 23 h, confirming that there is no appreciable homolytic pathway for H<sub>2</sub> generation (*SI Appendix*, Fig. S18).] Application of these guidelines to metal hydrides prone to homocoupling to form H<sub>2</sub> would be expected to impact the pH-dependent H<sub>2</sub> evolution activity, present challenges to isolating the metal hydride under electrolytic conditions without generating H<sub>2</sub>, and lead to lower overall selectivity.

The reduction potential required to generate  $[HPt(dmpe)_2]$ (PF<sub>6</sub>) (2) from the resting-state catalyst  $[Pt(dmpe)_2](PF_6)_2$  (3) (shown as  $E_{red1}$  in Eq. 7) is positive of the reduction potential for 2 ( $E_{red2}$  in Eq. 7):

$$\left[LM\right]^{n+1} \frac{2e^{-}, H^{+}}{E_{red1}} \left[LMH\right]^{n} \frac{e^{-}}{E_{red2}} \left[LMH\right]^{n-1}.$$
[7]

Facile electrolytic generation of  $[HPt(dmpe)_2](PF_6)$  (2) is possible, because  $E_{red1}$  is greater than  $E_{red2}$  (-1.7 vs. -2.8 V vs.  $Fe(C_5H_5)_2^{+/0}$ ) (shown in *SI Appendix*, Fig. S19). As a result, additional reduction of  $[HPt(dmpe)_2](PF_6)$  (2) to another intermediate is not possible at the electrochemical potential required to reduce **3**. Several other group 10 metal hydride complexes (26) along with multiple cobalt hydride complexes (84, 88, 92–94) have the same property; reduction of the hydride complex ( $[LMH]^n$ in Eq. **7**) is more challenging than reduction of the parent complex ( $[LM]^{n+1}$  in Eq. **7**) to generate the hydride (or  $E_{red1} > E_{red2}$ ). In cases where this property does not hold [i.e., the reduction potential of the hydride intermediate ( $E_{red2}$ ) is positive of the parent complex], it will likely be reduced under electrolytic conditions to generate a stronger hydride donor. For example, cobaloxime hydride



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complexes will generally be reduced further at potentials necessary to generate the hydride  $[LMH]^n$  in Eq. 7 (or  $E_{red2} > E_{red1}$ ). In these cases, the hydricity of the reduced complex ( $[LMH]^{n-1}$  in Eq. 7) can be applied to understand the  $pK_a$ -dependent hydrogen evolution catalysis, as  $[LMH]^{n-1}$  will likely be generated under electrolytic conditions.

### Conclusion

Despite immense interest in electrolytic fuel generation, there are few guidelines for the rational design of catalysts for selective  $CO_2$  reduction. We use thermodynamic relationships to understand the reactivity of metal hydrides, a branch point in the reactivity for formation of either H<sub>2</sub> or HCO<sub>2</sub><sup>-</sup>. We applied our analysis to construct a diagram that defines catalyst parameters for achieving selective  $CO_2$  reduction by targeting an appropriate hydricity. Hydricity is akin to an activity descriptor for H<sub>2</sub> and HCO<sub>2</sub><sup>-</sup> generation—a thermodynamic quantity that describes the bond energy of a key intermediate. Like an activity descriptor, hydricity is general across many classes of compounds, expanding its utility. Identifying descriptors in molecular systems is particularly advantageous, as bond energies are easily tuned through metal–ligand design.

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We applied our thermodynamic product diagram to identify a metal hydride with appropriate hydricity to reduce  $CO_2$  to formate under conditions that minimized concomitant  $H_2$  evolution. We expect that the thermodynamic framework described herein will continue to stimulate the discovery of more selective and efficient catalysts necessary for a carbon-neutral energy economy.

### Methods

Experimental details, cyclic voltammograms, NMR spectra, data from electrolysis, H<sub>2</sub> (GC) and HCO<sub>2</sub><sup>-</sup> (NMR) calibration curves, and additional thermodynamic product diagrams in DMSO and H<sub>2</sub>O are provided in *SI Appendix*. *SI Appendix* includes synthetic methods and materials, physical methods, electrochemical methods (including controlled potential electrolysis and product analysis), additional thermodynamic product diagrams, and NMR spectra.

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