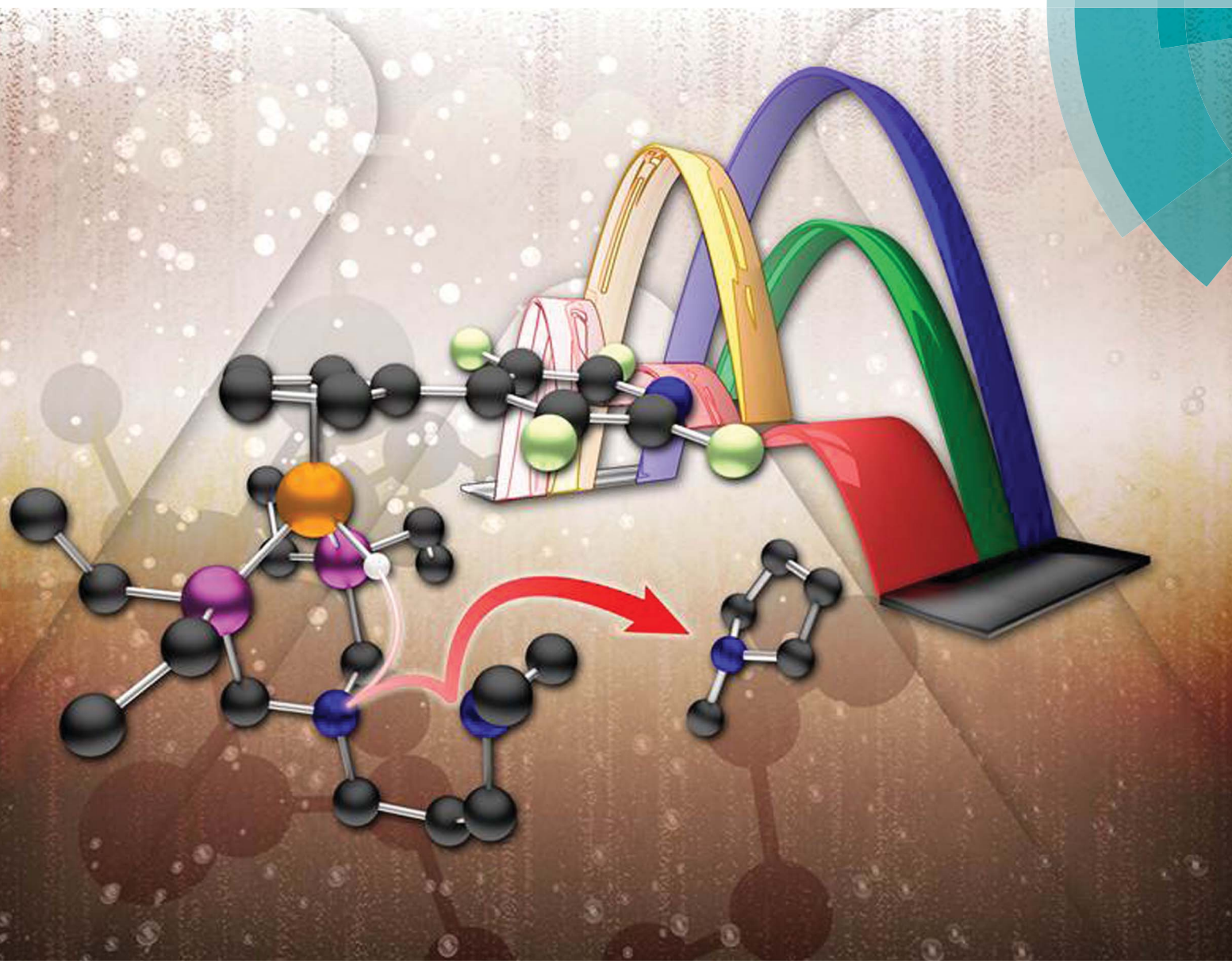


# Chemical Science

[www.rsc.org/chemicalscience](http://www.rsc.org/chemicalscience)



ISSN 2041-6539



ROYAL SOCIETY  
OF CHEMISTRY

EDGE ARTICLE

Monte L. Helm *et al.*

Increasing the rate of hydrogen oxidation without increasing the overpotential: a bio-inspired iron molecular electrocatalyst with an outer coordination sphere proton relay



# Increasing the rate of hydrogen oxidation without increasing the overpotential: a bio-inspired iron molecular electrocatalyst with an outer coordination sphere proton relay<sup>†</sup>

 Jonathan M. Darmon, Neeraj Kumar, Elliott B. Hulley, Charles J. Weiss,  
 Simone Raugei, R. Morris Bullock and Monte L. Helm\*

Oxidation of hydrogen ( $H_2$ ) to protons and electrons for energy production in fuel cells is currently catalyzed by platinum, but its low abundance and high cost present drawbacks to widespread adoption. Precisely controlled proton removal from the active site is critical in hydrogenase enzymes in nature that catalyze  $H_2$  oxidation using earth-abundant metals (iron and nickel). Here we report a synthetic iron complex,  $(Cp^{C_5F_5})Fe(P^{Et}_2N^{(CH_2)_3NMe_2P^{Et}_2})(Cl)$ , that serves as a precatalyst for the oxidation of  $H_2$ , with turnover frequencies of  $290\ s^{-1}$  in fluorobenzene, under 1 atm of  $H_2$  using 1,4-diazabicyclo[2.2.2]octane (DABCO) as the exogenous base. The inclusion of a properly tuned outer coordination sphere proton relay results in a cooperative effect between the primary, secondary and outer coordination spheres for moving protons, increasing the rate of  $H_2$  oxidation without increasing the overpotential when compared with the analogous complex featuring a single pendant base. This finding emphasizes the key role of pendant amines in mimicking the functionality of the proton pathway in the hydrogenase enzymes.

 Received 2nd February 2015  
 Accepted 4th March 2015

DOI: 10.1039/c5sc00398a

[www.rsc.org/chemicalscience](http://www.rsc.org/chemicalscience)

## Introduction

Nature has developed the ability to perform catalytic chemical transformations at amazing rates and efficiencies. For example, hydrogenase enzymes are able to both produce and oxidize  $H_2$  at rates exceeding  $10\ 000\ s^{-1}$  and at low overpotentials (high efficiencies).<sup>1–3</sup> Such remarkable performance has only been achieved through active sites that have evolved to minimize free energy differences between the reactive intermediates of the catalytic process, with the outer coordination sphere organized to shuttle protons between the surrounding matrix and the metal center.<sup>4</sup> Understanding and mimicking the function of hydrogenase enzymes is a particularly compelling challenge, as increasing efforts worldwide are devoted to utilizing clean, renewable resources for power. Solar and wind have the potential to eliminate our dependence on fossil fuels; however, their irregular geographic and temporal availability underlies the necessity

for efficient energy storage technologies. One promising strategy is to electrochemically convert energy into chemical bonds in the form of a fuel such as  $H_2$ , which can be stored, transported and used when needed.<sup>5</sup> Current  $H_2$  fuel cell technologies rely on high-cost, low abundant metals, such as platinum, to accomplish the oxidation of  $H_2$ , limiting their widespread use.<sup>6</sup> Hydrogenase enzymes, which use earth-abundant Ni and Fe metals, serve as inspiration to develop non-precious metal molecular catalysts for  $H_2$  production and oxidation with comparable rates and efficiencies.<sup>7–10</sup>

Development of functional mimics of the [FeFe]-hydrogenase has focused on optimizing the first and second coordination spheres of the active catalytic site.<sup>11–15</sup> Less attention has been paid towards the development and utility of the outer coordination sphere in synthetic catalysts to improve catalytic performance.<sup>16</sup> Although outer coordination sphere interactions may seem remote from the active site, recent work by Shaw has demonstrated that these effects can play a significant role in catalytic transformations.<sup>16</sup> Synthetic iron complexes that heterolytically cleave  $H_2$  have appeared in the literature including those that catalyze the oxidation of  $H_2$  using a chemical oxidant.<sup>17–19</sup> Our Center reported  $(Cp^{C_6F_5})Fe(P^{tBu}_2N^{Bn}_2)H$  ( $P^R_2N^{R'}_2 = RP(CH_2N(R')CH_2)_2PR$ ), a molecular iron electrocatalyst for the oxidation of  $H_2$  with an active site that has been thermodynamically tuned through modification of the first coordination sphere, and a pendant amine in the

Center for Molecular Electrocatalysis, Physical Sciences Division, Pacific Northwest National Laboratory, P.O. Box 999, K2-57, Richland, Washington 99352, USA.  
 E-mail: monte.helm@pnnl.gov

† Electronic supplementary information (ESI) available: Experimental procedures, syntheses, X-ray crystallographic data (CIF), and detailed theoretical discussion of presented and alternative mechanisms. CCDC 1046291. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c5sc00398a



second coordination sphere to heterolytically cleave  $\text{H}_2$ .<sup>20</sup> Very recently, we reported the related  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{R}}\text{N}^{\text{R}'}\text{P}^{\text{R}})\text{Cl}$  family of complexes ( $\text{P}^{\text{R}}\text{N}^{\text{R}'}\text{P}^{\text{R}} = \text{R}_2\text{PCH}_2\text{N}(\text{R}')\text{CH}_2\text{PR}_2$ ) for molecular electrocatalytic oxidation of  $\text{H}_2$  that included a detailed mechanistic study of the catalytic process.<sup>21</sup> In that study, deprotonation of the Fe(III) hydride was identified as having a significant thermodynamic barrier due to the mismatch between the  $\text{pK}_a$  values of the metal hydride and that of the protonated pendant base, limiting the rate of electrocatalytic  $\text{H}_2$  oxidation. We now report the synthesis of a modified iron complex that incorporates an *additional* proton relay in the *outer* coordination sphere to facilitate intramolecular proton movement. This new iron complex,  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})(\text{Cl})$ , **1-Cl**, electrocatalytically oxidizes  $\text{H}_2$  (1 atm) at a rate more than two orders of magnitude faster than the first iron-based, homogeneous  $\text{H}_2$  oxidation electrocatalyst,  $(\text{Cp}^{\text{C}_5\text{F}_5})\text{Fe}(\text{P}^{\text{tBu}}_2\text{N}^{\text{Bn}}_2)(\text{H})$  and one order of magnitude faster than the second generation  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{R}}\text{N}^{\text{R}'}\text{P}^{\text{R}})(\text{H})$  catalysts.<sup>20,21</sup> Equally remarkable, the addition of an outer sphere proton relay to a  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{R}}\text{N}^{\text{R}'}\text{P}^{\text{R}})(\text{H})$  catalyst increased the rate *without* an increase in the overpotential, illustrating the power of controlling proton movement for improved catalytic performance. Our study clearly demonstrates that outer coordination sphere effects can significantly enhance the performance of bio-inspired synthetic molecular catalysts.

## Results

### Synthesis and characterization of $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})(\text{Cl})$ (**1-Cl**) and electrocatalytic intermediates

The green iron chloride complex, **1-Cl**, was prepared by treating  $(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})\text{FeCl}_2$  with  $\text{NaCp}^{\text{C}_5\text{F}_4\text{N}}$ , and isolated in 92% yield. Reaction of **1-Cl** with  $\text{NaBAr}^{\text{F}}$  ( $\text{Ar}^{\text{F}} = 3,5\text{-bis}(\text{trifluoromethyl})\text{phenyl}$ ) under an atmosphere of  $\text{H}_2$  generated the protonated iron hydride species, **1-FeH(NH)**, which has been characterized by heteronuclear NMR spectroscopy and single crystal X-ray diffraction (Fig. 1, and ESI Table S1†). Spectroscopic NMR studies of **1-FeH(NH)** in solution were diagnostic of dynamic behavior involving the  $\text{H}_2$  ligand (see below); however, at  $-40^\circ\text{C}$ , the  $^1\text{H}$  NMR spectrum exhibits iron hydride ( $-17.9$  ppm) and protonated amine ( $10.4$  ppm) resonances, indicative of heterolytically cleaved  $\text{H}_2$ . In the solid state (Fig. 1), **1-FeH(NH)** adopts a three-legged piano-stool geometry, where the six-membered ring comprised of the PNP backbone and the iron center is in a boat conformation. The structure illustrates that heterolytic cleavage of  $\text{H}_2$  has occurred, as was also observed in the recently reported neutron diffraction structure of  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{H})(\text{P}^{\text{tBu}}_2\text{N}^{\text{Bn}}_2\text{H})$ .<sup>22</sup> The proton located on the outer coordination sphere amine forms a six-membered ring with the second coordination sphere pendant base through hydrogen bonding. Addition of five equivalents of

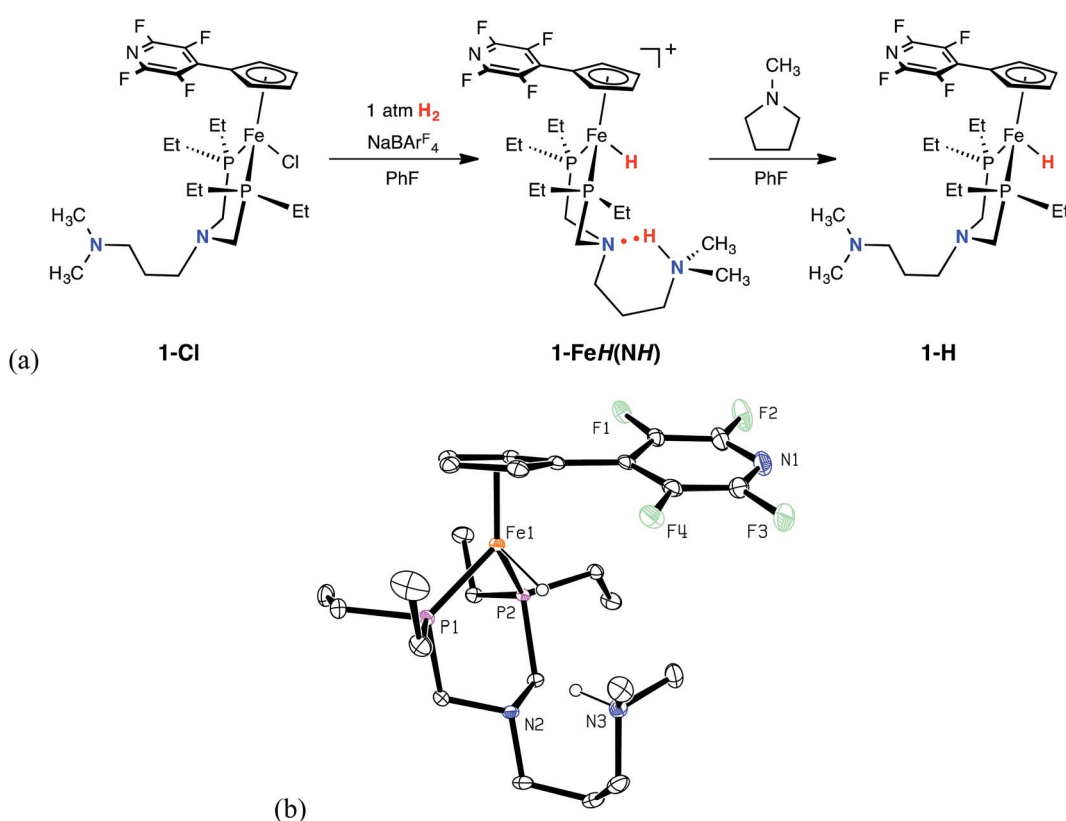


Fig. 1 (a) Observable catalytic intermediates from the reaction of **1-Cl** with  $\text{NaBAr}^{\text{F}}$  under  $1 \text{ atm H}_2$ . (b) Representation of the solid-state structure of **1-FeH(NH)** with 30% probability ellipsoids. The  $\text{BAr}^{\text{F}}$  anion and all hydrogen atoms with the exception of the proton and the hydride are omitted for clarity.



N-methylpyrrolidine deprotonated complex **1-FeH(NH)** and furnished the iron hydride species, **1-H**, as a viscous, red oil in 51% yield.

### Electrochemical characterization of the iron complexes and electrocatalytic oxidation of H<sub>2</sub>

Electrochemical characterization of the iron complexes were carried out in fluorobenzene solutions to avoid catalyst inhibition from solvent binding to the 16-electron complex.<sup>21</sup> The cyclic voltammogram (CV) of **1-Cl** exhibits a reversible one-electron oxidation wave assigned to the Fe<sup>II/III</sup> couple at  $-0.42$  V *versus* the ferrocenium/ferrocene couple (Fig. 2a). Irreversible oxidation waves for the protonated iron hydride, **1-FeH(NH)** ( $E_p = -0.36$  V) and the iron hydride, **1-H** ( $E_p = -0.61$  V) are observed in their CVs (Fig. 2a). Electrocatalytic oxidation studies of H<sub>2</sub> are carried out after addition of NaBAr<sub>4</sub><sup>F</sup> to **1-Cl** under 1 atm H<sub>2</sub> in fluorobenzene. Cyclic voltammograms recorded upon subsequent additions of either 1,4-diazabicyclo[2.2.2]octane (Fig. 2b, DABCO,  $pK_a = 18.2$  for H-DABCO<sup>+</sup> in CH<sub>3</sub>CN) or N-methylpyrrolidine (Fig. S1†,  $pK_a = 18.5$  for N-methylpyrrolidinium in CH<sub>3</sub>CN) result in a current enhancement consistent with electrocatalytic H<sub>2</sub> oxidation.<sup>23,24</sup> Base concentration and scan rate independence is achieved at 0.1 M DABCO and 6 V s<sup>-1</sup> (Fig. 2c and S2†), indicating that steady state conditions were achieved, yielding a reliable catalytic current ( $i_{cat}$ ) for calculation of the turnover frequency (TOF). Similar results were obtained when using N-methylpyrrolidine as exogenous base (Fig. S1†). Bulk electrolysis of a 0.5 mM solution of **1** in the presence of 1 atm of H<sub>2</sub> and 77 mM N-methylpyrrolidine confirmed H<sub>2</sub> oxidation through the quantitative analysis of the protons *via* NMR spectroscopy, resulting in an

average current efficiency of 95% and a minimum turnover number of 6.

$$\frac{i_{cat}}{i_p} = \frac{n}{0.4463} \sqrt{\frac{RTk_{obs}}{F\nu}} \quad (1)$$

The TOF for catalytic H<sub>2</sub> oxidation with **1-Cl** was calculated using eqn (1).<sup>25,26</sup> In this equation,  $i_p$  is the current under non-catalytic conditions,  $i_{cat}$  is the catalytic current,  $n$  is the number of electrons (2) involved in the catalytic reaction,  $k_{obs}$  is the observed first-order rate constant (TOF),  $R$  is the universal gas constant,  $T$  is the temperature in Kelvin,  $F$  is Faraday's constant,  $\nu$  is the scan rate, and 0.4463 is a constant determined by numerical solution of the diffusion equations.<sup>27</sup> The TOF for the oxidation of H<sub>2</sub> catalyzed by **1-Cl** was determined to be 290 s<sup>-1</sup> in the presence of  $\geq 77$  mM DABCO under 1 atm of H<sub>2</sub> at a half-wave potential ( $E_{cat/2}$ ) of  $-0.46$  V. The TOF for the oxidation of H<sub>2</sub> catalyzed by **1-Cl** was determined to be 180 s<sup>-1</sup> in the presence of  $\geq 65$  mM N-methylpyrrolidine, the base used in previously reported catalytic studies with (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(Cl) (Fig. S1†). The thermodynamic potential for the oxidation of H<sub>2</sub> using a 10 : 1 (base to acid) buffered solution of DABCO was determined by open circuit potential measurement and was found to be 1.22 V, resulting in a 0.65 V overpotential at  $E_{cat/2}$  (Fig. S3–S5†).<sup>28,29</sup> A buffered solution was used for the determination of the overpotential to minimize the experimental uncertainty in the measurement, in accord with recent recommendations.<sup>29</sup>

## Discussion

### Comparison of (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>(CH<sub>2</sub>)<sub>3</sub>Me</sup>P<sup>Et</sup>)(X), (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(X) and (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(ET<sub>2</sub>P(CH<sub>2</sub>)<sub>3</sub>PEt<sub>2</sub>)(X) (X = Cl, H)

To illustrate the impact of the number of proton relays on the rate of H<sub>2</sub> oxidation, electrochemical and electrocatalytic studies using iron complexes without pendant amines ((Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(depp)(X) (X = Cl, H), Scheme 1) and with one pendant amine ((Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(X), Scheme 1) were conducted under identical conditions to those of **1-X**. For the three Fe complexes, the reversible Fe(III/II) couple of the Cl<sup>-</sup> complexes differ by only 40 mV (Table 1 and Fig. S6†). Similarly, the peak of the waves of the Fe(II) hydride complexes range from  $-0.56$  to  $0.61$  V (Table 1 and Fig. S7†). The (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(depp)(H) (III/II) couple is reversible, while the (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(H) (III/II) wave is irreversible, but becomes reversible at scan rates  $\geq 0.5$  V s<sup>-1</sup>. The Fe(III/II) couple for **1-H** is irreversible up to scan rates as fast as 50 V s<sup>-1</sup> (Fig. S8†). The irreversible nature of the Fe(II) hydride oxidation of (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(H) and **1-H** can be attributed to the intramolecular deprotonation of the hydride by the pendant amines.<sup>21</sup> In the presence of exogenous base, the reversibility of the Fe(II) hydride oxidation waves as a function of scan rate can be used to estimate pseudo-first-order rate constants for intermolecular deprotonation. For (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(depp)(H) this rate was estimated to be  $<0.1$  s<sup>-1</sup> and  $>25$  s<sup>-1</sup> for (Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>)Fe(P<sup>Et</sup>N<sup>Me</sup>P<sup>Et</sup>)(H).<sup>21</sup> Similar studies of **1-H** show no reversibility, indicating the rate of intermolecular

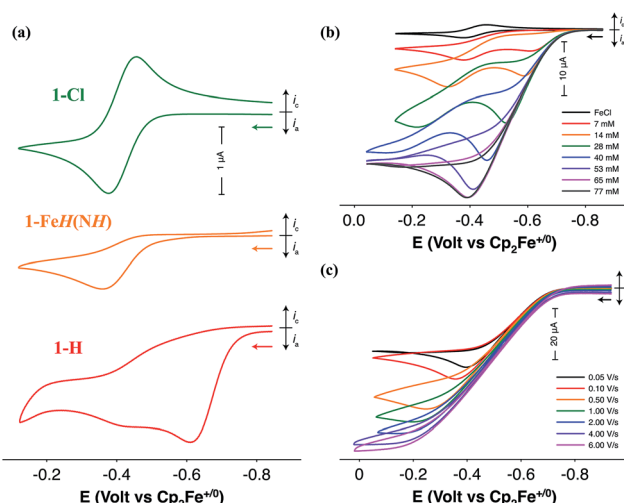
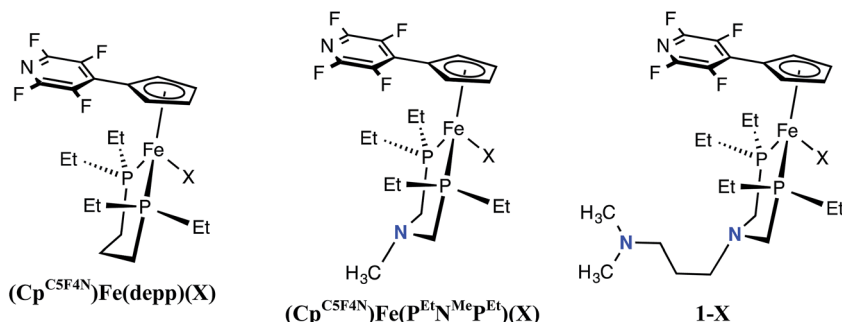


Fig. 2 (a) CVs of **1-Cl** (green), **1-FeH(NH)** (orange), and **1-H** (red) in fluorobenzene solution at  $50$  mV s<sup>-1</sup>. (b) CVs of a fluorobenzene solution of **1-FeH(NH)** upon addition of DABCO under 1 atm H<sub>2</sub> at  $50$  mV s<sup>-1</sup>. The initial CV (black) corresponds to **1-Cl** prior to activation with NaBAr<sub>4</sub><sup>F</sup>. (c) CVs of a fluorobenzene solution of **1-H** in the presence of 77 mM DABCO under 1 atm H<sub>2</sub> as a function of scan rate. Conditions for all experiments: 1 mM [Fe], 0.1 M [<sup>7</sup>Bu<sub>4</sub>N][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>], 22 °C.





Scheme 1 Complexes compared to examine the role of the number of pendant amines ( $X = \text{Cl}$ , or  $\text{H}$ ).

deprotonation is likely  $\gg 25 \text{ s}^{-1}$  (Fig. S9†). This data indicates that providing a pathway to move the proton further away from the metal center results in more facile deprotonation by the exogenous base.

Electrocatalytic  $\text{H}_2$  oxidation with **1-Cl** resulted in a TOF of  $180 \text{ s}^{-1}$  using *N*-methylpyrrolidine as the exogenous base and a TOF of  $290 \text{ s}^{-1}$  when using DABCO. Although *N*-methylpyrrolidine and DABCO have similar  $pK_a$  values, the less sterically demanding nature of DABCO yields faster rates of  $\text{H}_2$  oxidation, indicating deprotonation by exogenous base is likely a rate-limiting factor. In order to study the effect of the outer coordination sphere pendant amine on catalysis,  $\text{H}_2$  oxidation TOFs and overpotentials with  $\text{Fe}(\text{depp})(\text{X})$  and  $\text{Fe}(\text{PEtN}^{\text{Me}}\text{PEt})(\text{X})$  and DABCO were determined (Table 1 and Fig. S10 and 11†). The rates of  $\text{H}_2$  oxidation by the three compounds increase as the number of pendant amines increases, with no significant change ( $<10 \text{ mV}$ ) in the overpotential. This data is again consistent with the hypothesis that increasing the rate of deprotonation of the  $\text{Fe}(\text{III})$  hydride increases the rate of catalysis, rather than an increase attributable to the driving force (overpotential) of catalysis.

### Proposed mechanism of electrocatalytic $\text{H}_2$ oxidation

The proposed mechanism for  $\text{H}_2$  oxidation by **1-Cl** is shown in Fig. 3 (some intermediates have been omitted for clarity; see Fig. S12–14† for complete mechanistic details). Addition of  $\text{H}_2$  to the unsaturated, 16-electron  $\text{Fe}(\text{II})$  complex is followed by heterolytic cleavage to form the protonated iron hydride **1-FeH(NH)**. A dihydrogen bonding<sup>30</sup> interaction between the

proton and hydride in this complex is likely, as previously observed in the analogous  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{H})(\text{P}^{\text{tBu}}_2\text{N}^{\text{tBu}}_2\text{H})]^+$  complex.<sup>22</sup> Computed thermodynamic and kinetic parameters for these steps (Fig. 3, steps 1–3) suggest that the resulting species are close in free energy and the barriers are less than 9 kcal mol<sup>−1</sup> (Fig. S13†), and therefore do not limit the observed rate of oxidation of  $\text{H}_2$ . These results are further supported by the dynamic behavior observed by <sup>1</sup>H NMR spectroscopy (Fig. S15†). At  $-40^\circ\text{C}$  (at 500 MHz), the proton and hydride environments of **1-FeH(NH)** are resolved, and the  $\text{FeH}$  ( $-17.93 \text{ ppm}$ ) and  $\text{NH}$  ( $10.39 \text{ ppm}$ ) resonances are independently observed. This peak separation corresponds to a proton hydride exchange of  $\sim 6.4 \times 10^3 \text{ s}^{-1}$  ( $\Delta G^\ddagger$  of 12 kcal mol<sup>−1</sup> according to simple transition state theory arguments) at the coalescence temperature of  $\sim 20^\circ\text{C}$ . The coalesced proton-hydride resonance is observed at  $-6.1 \text{ ppm}$  ( $65^\circ\text{C}$ ). This deviates from the expected value of about  $-3.8 \text{ ppm}$ , and is consistent with contributions from other energetically-matched species (*i.e.*  $\text{Fe}(\eta^2\text{-H}_2)$  complex). In addition to the fluxional behavior of the proton and hydride resonances, the ethyl and methylene resonances of  $\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{N}^{\text{Me}}}\text{P}^{\text{Et}}$  also coalesce at elevated temperatures (Fig. S15†). At  $-40^\circ\text{C}$  (500 MHz), the two ethyl environments are resolved, and the two resonances are independently observed (0.71 and 0.81 ppm). This peak separation corresponds to a Fe-PNP inversion (Fig. S16†) rate of  $25 \text{ s}^{-1}$  at the coalescence temperature of  $\sim 40^\circ\text{C}$  ( $\Delta G^\ddagger$  of 16 kcal mol<sup>−1</sup>). This behavior is consistent with loss of  $\text{H}_2$  and formation of the pseudosymmetric, fluxional 16-electron species, and is consistent with the computational results. Given the calculated energies and this experimentally determined rate for heterolytic

Table 1 Electrochemical and electrocatalytic data for complexes with increasing number of proton relays

Compound	Fe(III/II) couple <sup>a</sup> (V) $X = \text{Cl}$	Fe(III/II) couple <sup>b</sup> (V) $X = \text{H}$	TOF <sup>c</sup> ( $\text{s}^{-1}$ )	Overpotential <sup>d</sup> (V)
$(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{depp})(\text{X})$	−0.45	−0.60	NC <sup>e</sup>	—
$(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEtN}^{\text{Me}}\text{PEt})(\text{X})$	−0.41	−0.56	34	0.64
<b>1-X</b>	−0.42	−0.61	290	0.65

<sup>a</sup> Conditions: fluorobenzene solution, 1 mM [Fe], 0.1 M [<sup>n</sup>Bu<sub>4</sub>N][B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub>], 50 mV s<sup>−1</sup>, 22 °C. <sup>b</sup> Values reported are for the oxidative peak potential.

<sup>c</sup> Reported values were determined upon addition of a DABCO solution, and are given under base- and scan rate-independent conditions (see above and the ESI† for additional information). <sup>d</sup> Reported values were determined upon addition of a 10 : 1 DABCO to DABCO-H<sup>+</sup> solution as the exogenous base, and are given under base- and scan rate-independent conditions (see above and the ESI† for additional information). <sup>e</sup> Not Catalytic (NC) – no significant current enhancement was observed, indicating this complex is not a catalyst for  $\text{H}_2$  oxidation.

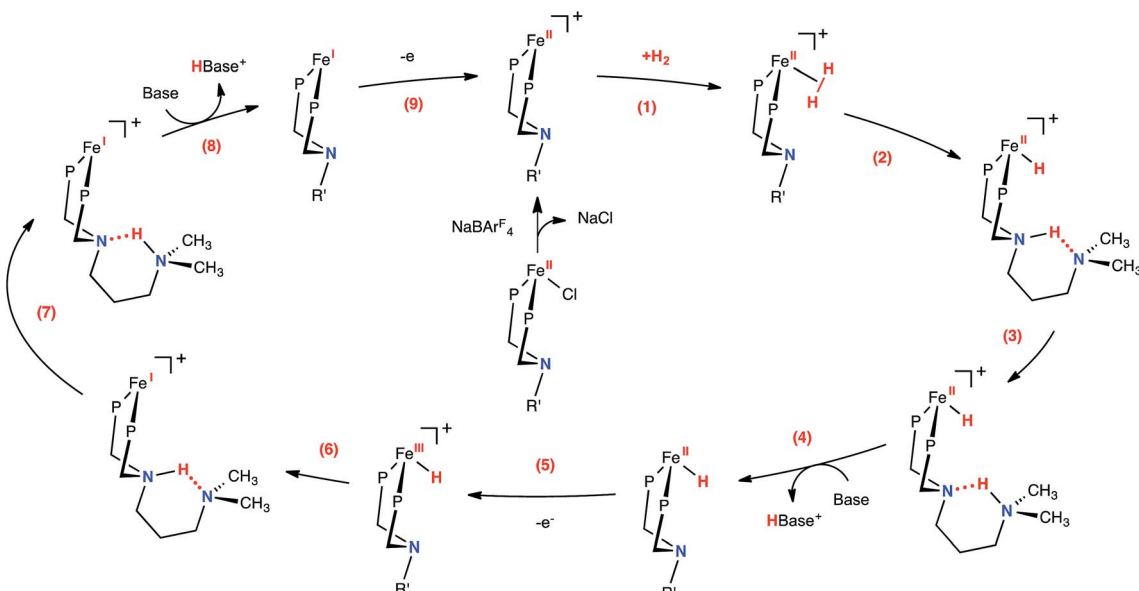


Fig. 3 Proposed mechanism for the electrocatalytic oxidation of H<sub>2</sub> by  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEtN}^{(\text{CH}_2)_3\text{NMe}_2}\text{PEt})]^{+}$ . Steps (4) and (8) include opening of the six-membered ring (not shown), prior to deprotonation by exogenous base (see Fig. 4, S13 and S14†). The Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup> and ethyl substituents on the phosphorus atoms have been omitted for clarity (R' = CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>N(CH<sub>3</sub>)<sub>2</sub>).

cleavage, it is apparent that these steps are not rate-limiting for catalysis in this system.

Consistent with our current study, previous experimental and computational work describing the mechanism of H<sub>2</sub> oxidation by the Fe-P<sup>R</sup>N<sup>R'</sup>P<sup>R</sup> family of complexes indicated that H<sub>2</sub> addition and cleavage (*i.e.*, Fig. 3, steps 1 and 2) and the subsequent deprotonation of the pendant amine of the Fe(<sup>II</sup>) hydride (*i.e.*, Fig. 3, steps 3–4) do not significantly contribute to

the overall catalytic rate. In contrast, the intramolecular followed by intermolecular deprotonation steps of the Fe(<sup>III</sup>) hydride are the steps that limit the overall observed rate of catalysis (Fig. 3, steps 6–8, Fig. 4).<sup>21</sup> The large difference in acidity between the Fe(<sup>III</sup>) hydride and the protonated, secondary sphere pendant amine (9 pK<sub>a</sub> units) makes the intramolecular proton transfer thermodynamically unfavorable. Additionally, the poor steric accessibility of the

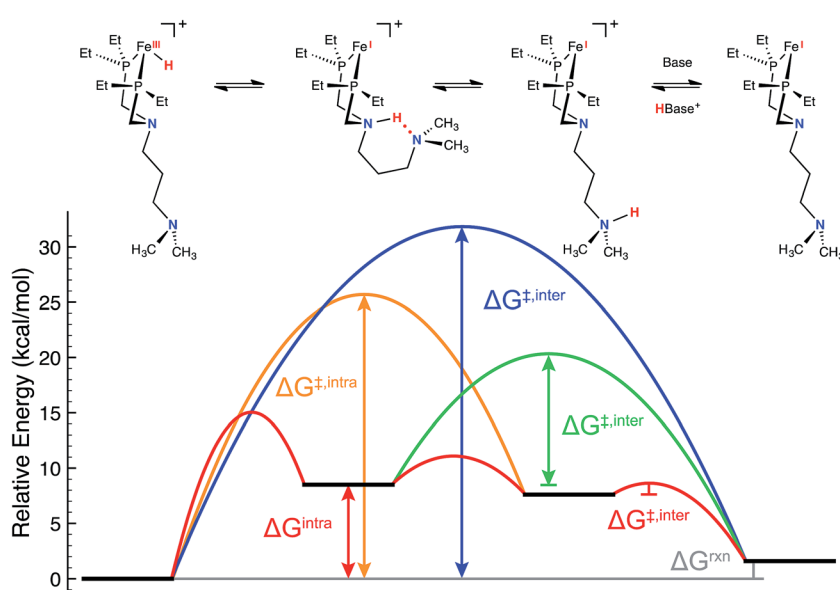


Fig. 4 Free energy diagram for the key intra- and intermolecular deprotonation steps of  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEtN}^{(\text{CH}_2)_3\text{NMe}_2}\text{PEt})(\text{H})]^{+}$  by N-methylpyrrolidine as obtained from density functional theory calculations (see ESI† for additional details). Red lines: lowest-free energy deprotonation pathway (upper reaction scheme); blue line: direct deprotonation of the iron(<sup>III</sup>) hydride by the exogenous base; orange line: deprotonation of the iron(<sup>III</sup>) hydride by the outer coordination sphere pendant amine; green line: deprotonation of the second coordination sphere pendant amine by exogenous base. The Cp<sup>C<sub>5</sub>F<sub>4</sub>N</sup> substituent has been omitted for clarity.

protonated pendant amine introduces a further penalty for removal of the proton by an exogenous base (*e.g.* Fig. 4, green  $\Delta G_{\text{inter}}^{\ddagger}$ ). A detailed computational analysis of  $\text{H}_2$  oxidation by  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEt}_2\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})]^+$  using *N*-methylpyrrolidine as the exogenous base was carried out, allowing for direct comparison to the previously published  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEt}_2\text{N}^{\text{Me}}\text{P}^{\text{Et}})]^+$  complex.<sup>21</sup> The calculations revealed that introduction of an outer coordination sphere amine results in a more suitably energy-matched pathway for the proton from the Fe(III) center to the periphery of the catalyst (Fig. 4, red trace). The presence of the outer coordination sphere amine stabilizes the aminium in the second coordination sphere through intramolecular hydrogen bonding, which results in a reduced  $\text{p}K_{\text{a}}$  difference between the Fe(III) hydride intermediate and protonated amine by about 3  $\text{p}K_{\text{a}}$  units with respect to the  $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PEt}_2\text{N}^{\text{Me}}\text{P}^{\text{Et}})]^+$  complex (Fig. S9†).<sup>21</sup> Proton transfer between the two pendant amines is very facile, and the intramolecular hydrogen bond between them easily breaks, exposing the protonated outer sphere amine to the exogenous base, diminishing the steric penalty for deprotonation. It is important to point out that direct deprotonation of the Fe(III) hydride either by the outer sphere amine (Fig. 4, orange  $\Delta G_{\text{intra}}^{\ddagger}$ ) or by an exogenous base (Fig. 4, blue  $\Delta G_{\text{inter}}^{\ddagger}$ ) is kinetically disfavored because of the scarce accessibility of the base to the metal hydride. A detailed mechanistic discussion of the various pathways for the proton movement is provided in the ESI.† As observed in the remarkable TOF exhibited by this catalyst, the low energy pathway provided by the addition of the outer coordination sphere pendant amine to move the proton away from the Fe(III) active site mimics the functionality of hydrogenase enzymes, in which protons shuttle through different amino acids for rapid proton movement.

## Conclusions

These results conclusively demonstrate that controlled movement of protons from the primary, to the secondary and outer coordination spheres dramatically improves the rate of  $\text{H}_2$  oxidation in an iron-based molecular electrocatalyst without increasing the overpotential. Although this report focuses on a single catalyst, the comprehensive thermodynamic analysis highlights the broader implications of this approach, specifically how mimicking enzyme functionality to minimize free energy differences between catalytic intermediates can be used to improve catalyst performance. One of the most exciting results from this work is the demonstrated use of rational catalyst design to control proton movement and lower the barrier of the rate-limiting step in catalysis by the previously reported  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{PR}^{\text{R}}\text{N}^{\text{R}}\text{P}^{\text{R}})(\text{H})$  family. As electrocatalytic rates of  $\text{H}_2$  oxidation escalate, an increased focus on lowering the overpotential of these catalysts will be crucial.

## Experimental

All manipulations of free phosphine ligands and metal reagents were carried out under argon using standard vacuum line, Schlenk, and inert atmosphere glovebox techniques. Solvents

were purified by passage through neutral alumina using an Innovative Technology, Inc., PureSolv™ solvent purification system. THF-*d*<sub>8</sub> and bromobenzene-*d*<sub>5</sub> were vacuum transferred from NaK and CaH<sub>2</sub>, respectively after stirring for a minimum of 24 h under an inert atmosphere prior to use. DABCO was dried under vacuum prior to use. *N*-Methylpyrrolidine was dried over KOH, degassed, distilled under vacuum, and stored in an argon-filled glovebox. Elemental analyses were performed by Atlantic Microlab, Inc. in Norcross, GA. The following compounds were prepared according to literature procedures: NaCp<sup>C<sub>5</sub>F<sub>4</sub>N</sup>,<sup>22</sup> and  $[(\text{Et}_2\text{O})_2\text{H}][\text{B}(\text{C}_6\text{F}_5)_4]$ .<sup>31</sup> Additional experimental details can be found in the ESI, pages S3–S7.†

## Instrumentation

**NMR.** Spectra were recorded on a Varian Inova spectrometer (500 MHz for <sup>1</sup>H) at 22 °C unless otherwise noted. All <sup>1</sup>H chemical shifts have been internally calibrated using the monoprotio impurity of the deuterated solvent. The <sup>31</sup>P{<sup>1</sup>H} NMR spectra were referenced to external phosphoric acid at 0 ppm. The <sup>19</sup>F NMR spectra were referenced to external fluorobenzene at -113.15 ppm. A fluorobenzene solution of tetramethylsilane (3.0 mM) was used as the solvent for variable temperature NMR experiments.

**Electrochemistry.** All experimental procedures were conducted at ambient temperature, 22–23 °C, under argon in a vacuum atmospheres drybox. A standard three-electrode configuration was employed in conjunction with a CH Instruments potentiostat interfaced to a computer with CH Instruments 700 D software. All voltammetric scans were recorded using glassy-carbon working electrode disks of 1 mm diameter (Cypress Systems EE040). A glassy-carbon rod (Structure Probe, Inc.) and platinum wire (Alfa-Aesar) were used as auxiliary electrodes and quasi-reference electrodes, respectively. All glassware for electrochemical experiments was oven dried overnight and allowed to cool to room temperature before use.  $[\text{Cp}_2\text{Co}][\text{PF}_6]$  and  $(\eta^6\text{C}_6\text{H}_6)_2\text{Cr}$  were used as an internal standard, and all potentials reported within this work are measured *vs.* the  $\text{Cp}_2\text{Co}^{+/-}$  couple at -1.33 V or the  $(\eta^6\text{C}_6\text{H}_6)_2\text{Cr}^{+/-}$  couple at -1.22 V and are reported *vs.*  $\text{Cp}_2\text{Fe}^{+/-}$ . Bases were measured and transferred to electrochemical solutions *via* gas-tight syringes. Controlled potential electrolyses were performed using a high-power BASi potentiostat (Fig. S17†).

**UV-vis.** UV-vis data were collected under argon with an Ocean Optics USB2000+ spectrometer (Fig. S18†).

**X-ray diffraction.** The crystal selected for diffraction studies was immersed in Paratone-N oil, placed on a Nylon loop, and transferred to a precooled cold stream of N<sub>2</sub>. A Bruker KAPPA APEX II CCD diffractometer with 0.71073 Å Mo K $\alpha$  radiation was used. The space group was determined on the basis of systematic absences and intensity statistics. The structure was solved by direct methods and refined by full-matrix least squares on  $F^2$ . All non-hydrogen atoms were refined anisotropically unless otherwise stated. Hydrogen atoms were placed at idealized positions and refined using the riding model unless otherwise stated. Data collection and cell refinement were performed using Bruker APEX2 software. Data reduction and absorption

correction was performed using Bruker's SAINT and SADABS programs, respectively.<sup>32,33</sup> Structural solution and refinements were completed using SHELXS-97 and SHELXL-97 (ref. 34) using the OLEX2 software package<sup>35</sup> as a front-end.

**Computational details.** All possible reaction intermediates involved in the catalytic cycle and the transitions states between them were optimized *via* density function theory (DFT) calculations. The hybrid B3P86 functional<sup>36,37</sup> was used along with the Stuttgart–Dresden relativistic effective core potential for Fe and its associated basis set (SDD)<sup>38</sup> and the 6-31G\* basis set for all the atoms with one additional *p* polarization function<sup>39</sup> for the protic hydrogens. This level of theory has previously shown great accuracy for thermodynamic and kinetics parameters of first row transition metal based catalytic systems.<sup>40–42</sup> Harmonic vibrational frequencies were calculated at the optimized geometries using the same level of theory to estimate the zero-point energy (ZPE) and the thermal contributions (298 K and 1 atm) to the free energy. Transition states were verified by intrinsic reaction coordinate (IRC) calculations. All the molecular structures were optimized in fluorobenzene using the SMD continuum model<sup>43</sup> of solvation. Calculations were performed using *Gaussian* 09 suite of programs.<sup>44</sup>

### Preparation of $\text{Et}_2\text{PCH}_2\text{N}((\text{CH}_2)_3\text{NMe}_2)\text{CH}_2\text{PEt}_2$

A 50 mL Schlenk flask was loaded with 0.67 g (22 mmol) of paraformaldehyde, 15 mL of absolute ethanol, and a stir bar. Cycling between vacuum and nitrogen degassed the vessel, and 2.54 mL (22 mmol) of diethylphosphine was added by syringe with rapid stirring. After 0.5 h, 1.40 mL (11 mmol) of 3-(dimethylamino)-1-propylamine was added by syringe. The suspension rapidly became a clear, colorless solution over the course of 1 h. After 24 h, the solvent was removed under vacuum, and 3.20 g (94% yield) of a colorless oil identified as  $\text{Et}_2\text{PCH}_2\text{N}((\text{CH}_2)_3\text{NMe}_2)\text{CH}_2\text{PEt}_2$  was collected.  $^1\text{H}\{^3\text{P}\}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  1.06 (t, *J* = 7.6 Hz, 12H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.28–1.48 (m, 8H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.58 (dt, *J* = 14.2, 7.1 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.13 (s, 6H, N(CH<sub>3</sub>)<sub>2</sub>), 2.22 (app. t, *J* = 7.1 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.67 (app. t, *J* = 7.1 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.74 (s, 4H, PCH<sub>2</sub>N).  $^{31}\text{P}\{^1\text{H}\}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  –32.9 (s).  $^{13}\text{C}\{^1\text{H}\}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  15.5 (d, *J*<sub>CP</sub> = 14 Hz, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 19.6 (d, *J*<sub>CP</sub> = 13 Hz, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 26.5 (s, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 46.0 (s, N(CH<sub>3</sub>)<sub>2</sub>), 55.9 (t, *J*<sub>CP</sub> = 8 Hz, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 58.4 (dd, *J*<sub>CP</sub> = 5, 8 Hz, PCH<sub>2</sub>N), 58.7 (s, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>).

### Preparation of $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})(\text{Cl})$ (1-Cl)

A 20 mL scintillation vial was charged with 0.083 g (0.65 mmol) of FeCl<sub>2</sub>, 2 mL THF and a stir bar. A solution of  $\text{Et}_2\text{PCH}_2\text{N}((\text{CH}_2)_3\text{NMe}_2)\text{CH}_2\text{PEt}_2$  (0.20 g, 0.65 mmol) in 3 mL of THF was added to the suspension with rapid stirring, and the mixture rapidly became golden yellow. After stirring for 1 h, the resulting solution was stored at –35 °C for 30 minutes. A chilled (10 mL, –35 °C for 20 minutes) THF solution of NaCp<sup>C<sub>5</sub>F<sub>4</sub>N</sup> (0.16 g, 0.65 mmol) was added in a dropwise fashion with rapid stirring, causing an immediate color change to an intense green. After 24 h, the solvent was removed under vacuum. The

residue was extracted with diethyl ether, and the resulting mixture was filtered through a pad of Celite. The solvent was removed under vacuum to afford 0.37 g (92% yield) of a dark green solid identified as  $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})(\text{Cl})$ . The product was recrystallized from a saturated solution of diethyl ether at –35 °C to afford analytically pure material. Anal. calcd for C<sub>25</sub>H<sub>40</sub>ClF<sub>4</sub>FeN<sub>3</sub>P<sub>2</sub>: C, 49.08; H, 6.59; N, 6.87. Found: C, 49.34; H, 6.64; N, 6.89.  $^1\text{H}\{^3\text{P}\}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  1.19 (t, *J* = 7.7 Hz, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.23 (t, *J* = 7.7 Hz, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.55 (p, *J* = 7.0 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 1.94–2.08 (m, 4H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 2.11 (s, 6H, N(CH<sub>3</sub>)<sub>2</sub>), 2.14–2.38 (m, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub> and NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.26 (d, *J* = 12.6 Hz, 2H, NCH<sub>2</sub>P), 2.42 (t, *J* = 7.0 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.77 (d, *J* = 12.6 Hz, 2H, NCH<sub>2</sub>P), 3.77 (app. t, *J* = 2.1 Hz, 2H, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), 5.43 (app. p, *J* = 1.8 Hz, 2H, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N).  $^{31}\text{P}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  44.5 (s).  $^{19}\text{F}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  –141.0 (m), –96.6 (m).

### Characterization of $[(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{N}(\text{H})\text{Me}_2}\text{P}^{\text{Et}})(\text{H})]$ [BAr<sub>4</sub><sup>F</sup>] (1-FeH(NH))

A 5 mL scintillation vial was charged with **1** (0.015 g, 24  $\mu$ mol), NaBAr<sub>4</sub><sup>F</sup> (0.022 g, 24  $\mu$ mol), and a stir bar. Bromobenzene-*d*<sub>5</sub> (1 mL) was added with rapid stirring under an atmosphere of H<sub>2</sub>, resulting in a rapid color change from intensely green to a transparent red-orange. The mixture was filtered through a pad of Celite, and the resulting solution was characterized by NMR spectroscopy.  $^1\text{H}$  NMR (fluorobenzene, 25 °C):  $\delta$  0.75 (br m, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 0.86 (br m, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.18–1.37 (br m, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub> and NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 1.48 (br m, 4H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.67 (br d, *J* = 11.7 Hz, NCH<sub>2</sub>P), 2.19 (br d, *J* = 11.7 Hz, NCH<sub>2</sub>P), 2.27 (s, 6H, N(CH<sub>3</sub>)<sub>2</sub>), 2.31 (br t, *J* = 5 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.46 (br t, *J* = 5 Hz, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 4.40 (br s, 2H, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), 5.07 (br s, 2H, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), 7.63 (br s, 4H, BAr<sub>4</sub><sup>F</sup>), 8.29 (br s, 8H, BAr<sub>4</sub><sup>F</sup>).  $^{31}\text{P}$  NMR (bromobenzene-*d*<sub>5</sub>, 22 °C):  $\delta$  69.0 (br s).  $^{19}\text{F}$  NMR (fluorobenzene, 22 °C):  $\delta$  –143.2 (br s, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), –93.7 (br s, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), –62.3 (br s, BAr<sub>4</sub><sup>F</sup>).

### Preparation of $(\text{Cp}^{\text{C}_5\text{F}_4\text{N}})\text{Fe}(\text{P}^{\text{Et}}\text{N}^{(\text{CH}_2)_3\text{NMe}_2}\text{P}^{\text{Et}})(\text{H})$ (1-H)

A 20 mL scintillation vial was charged with **1** (0.15 g, 0.24 mmol), NaBAr<sub>4</sub><sup>F</sup> (0.22 g, 0.24 mmol), and a stir bar. Fluorobenzene (10 mL) was added with rapid stirring under an atmosphere of H<sub>2</sub>, resulting in a rapid color change from intensely green to a transparent red-orange. After 1 h, *N*-methylpyrrolidine (0.13 mL, 1.2 mmol) was added by micro-syringe and the solution immediately became purple-red in color. After stirring for 24 h, the solvent was removed under vacuum. The residue was extracted with toluene, and filtered through a pad of Celite. Removal of solvent afforded a viscous red oil (0.072 g, 51% yield) identified as **1-H**.  $^1\text{H}\{^3\text{P}\}$  NMR (THF-*d*<sub>8</sub>, 22 °C):  $\delta$  –17.0 (s, 1H, FeH), 0.93 (t, *J* = 7.6 Hz, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.12 (t, *J* = 7.6 Hz, 6H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.29–1.70 (m, 8H, P(CH<sub>2</sub>CH<sub>3</sub>)<sub>2</sub>), 1.69–1.87 (m, 4H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub> and PCH<sub>2</sub>N), 2.12 (s, 6H, N(CH<sub>3</sub>)<sub>2</sub>), 2.19 (t, *J* = 6.9, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub> or NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.34 (t, *J* = 7.0, 2H, NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub> or NCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NMe<sub>2</sub>), 2.88 (d, *J* = 12.0, 2H, PCH<sub>2</sub>N), 4.54 (app. t, *J* = 2.0 Hz, 2H, C<sub>5</sub>H<sub>4</sub>C<sub>5</sub>F<sub>4</sub>N), 5.04 (app. p, *J*



$\delta = 2.1$  Hz, 2H,  $C_5H_4C_5F_4N$ ).  $^{31}P$  NMR (THF- $d_8$ , 22 °C):  $\delta$  65.2 (d,  $J_{PH} = 71.9$  Hz).  $^{19}F$  NMR (THF- $d_8$ , 22 °C):  $\delta$  -144.7 (m), -96.8 (m).

### Preparation of $[(DABCO)_2H][B(C_6F_5)_4]$

A 20 mL scintillation vial was charged with DABCO (0.50 g, 4.4 mmol), 5 mL of fluorobenzene, and a stir bar. A fluorobenzene solution (10 mL) of  $[(Et_2O)_2H][B(C_6F_5)_4]$  (1.85 g, 2.23 mmol) was added dropwise, and resulted in the precipitation of a white solid. After 4 h, the solvent was removed, and a white solid (1.88 g, 93% yield) identified as  $[(DABCO)_2H][B(C_6F_5)_4]$  was isolated.  $^1H$  NMR (dichloromethane- $d_2$ , 22 °C):  $\delta$  2.91 (s, 24H,  $CH_2$ ), 13.81 (s, 1H, N-H).  $^{13}C\{^1H\}$  NMR (dichloromethane- $d_2$ , 22 °C):  $\delta$  46.0 (s).  $^{19}F$  NMR (dichloromethane- $d_2$ , 22 °C):  $\delta$  -166.7 (t,  $J = 18$  Hz,  $B(C_6F_5)_4$ ), -162.7 (t,  $J = 20$  Hz,  $B(C_6F_5)_4$ ), -132.4 (d,  $J = 10$  Hz,  $B(C_6F_5)_4$ ).

### Electrochemical $H_2$ oxidation catalyzed by $[(Cp^{C_5F_4N})Fe(P^{Et}N^{(CH_2)_3NMe_2}P^{Et})]$

A 1.0 mL fluorobenzene solution containing  $(Cp^{C_5F_4N})Fe(P^{Et}N^{(CH_2)_3NMe_2}P^{Et})(Cl)$  (1.0 mM) and  $[Bu_4N][B(C_6F_5)_4]$  (0.1 M) was prepared. Cyclic voltammograms were recorded at varying scan rates to determine the non-catalytic current ( $i_p$ ).  $NaBAr^{F_4}$  (0.9 mg) was added with rapid stirring under an atmosphere of  $H_2$  resulting in a color change from green to orange. Aliquots of a stock solution of DABCO (0.48 M) in fluorobenzene or a freshly prepared buffered solution of DABCO (see below) were added by microsyringe and cyclic voltammograms were collected after each addition.

### Bulk electrolysis

Bulk electrolysis was performed using a high-power BASi potentiostat with a four-necked flask. One neck was sealed with a rubber septum through which a copper wire was fed that pierced through a cylinder of reticulated vitreous carbon, and functioned as the working electrode. The second and third necks were equipped with a reference electrode and a counter electrode suspended from copper wire that was fed through rubber septa. A AgCl-coated silver wire suspended in a 0.1 M fluorobenzene solution of  $[Bu_4N][B(C_6F_5)_4]$  in a glass tube with a Vycor frit served as the reference electrode. The counter electrode consisted of a Ni-Cr coiled wire and a 0.1 M fluorobenzene solution of  $[Bu_4N][B(C_6F_5)_4]$  in a glass tube with a glass frit. The fourth neck was fitted with a septum through which a copper wire attached to a second working electrode (1 mm PEEK-encased glassy carbon, Cypress Systems EE040) was fed. The second working electrode was used to record cyclic voltammograms. A needle was placed through this septum to introduce  $H_2$  into the cell. The flask contained 10 mL of a 0.1 M  $[Bu_4N][B(C_6F_5)_4]$  solution in fluorobenzene as well as 1,  $(Cp^{C_5F_4N})Fe(P^{Et}N^{(CH_2)_3NMe_2}P^{Et})(Cl)$  (0.5 mM) and a small amount of  $Cp_2CoPF_6$  (~0.9 mM) as an internal reference. Before and after adding  $NaBAr^{F_4}$  (10 mg, 11  $\mu$ mol) and *N*-methylpyrrolidine (80  $\mu$ L, 77 mM), two cyclic voltammograms were recorded to ensure that the solution was under catalytic conditions and to determine the applied potential for bulk electrolysis.

Controlled-potential electrolysis was performed at approximately 0.1 V positive of the peak potential for  $(Cp^{C_5F_4N})Fe(P^{Et}N^{(CH_2)_3NMe_2}P^{Et})(H)$ . After the application of 4.01 C of charge (corresponding to 4.2 calculated turnovers) a 200  $\mu$ L aliquot of the solution was removed and mixed with 400  $\mu$ L of a 20 mM solution of Verkade's base (2,8,9-triisopropyl-2,5,8,9-tetraaza-1-phosphabicyclo[3.3.3]undecane, 30 mg dissolved in 5 mL tetrahydrofuran). Verkade's base ( $pK_a$  of the conjugate acid = 33.6 in  $CH_3CN$ )<sup>45</sup> sequestered all protons from the protonated *N*-methylpyrrolidine ( $pK_a$  of the conjugate acid = 18.5 in  $CH_3CN$ ) produced from bulk electrolysis, forming protonated Verkade's base. The amount of Verkade's base ( $\delta$  118.0) and protonated Verkade's base ( $\delta$  -11.9) can be quantified by  $^{31}P\{^1H\}$  NMR spectroscopy, based on the known concentration of the base. The amount of protons generated (3.6 mM in a 10 mL solution, 0.036 mmol, corresponding to 3.6 turnovers) from the oxidation of  $H_2$  can be determined. A second sample was taken after passing 5.90 C of charge (corresponding to 6.2 calculated turnovers), giving the amount of protons as 6.5 mM in a 9.8 mL solution (0.064 mmol, corresponding to 6.6 turnovers). An average Faradic efficiency of 95% was calculated for  $H_2$  oxidation.

### Open circuit potential measurement of a 10 : 1 DABCO to DABCO- $H^+$ solution

A stock solution of DABCO (0.030 g, 0.27 mmol),  $[(DABCO)_2H][B(C_6F_5)_4]$  (0.027 g, 30  $\mu$ mol) and  $[Bu_4N][B(C_6F_5)_4]$  (0.276 g, 0.30 mmol) in fluorobenzene was prepared by transferring the compounds to a 3 mL volumetric flask and diluting with solvent. A 1 mL aliquot of the solution was transferred to the electrochemical cell that was prepared according the method reported by Roberts and Bullock.<sup>28</sup> The solution was continuously sparged with  $H_2$ , and after 1 min, the open circuit potential was measured. Measurements were repeated until a consistent value was obtained, which signaled saturation of  $H_2$ . Following one more repeated open circuit potential measurement, an average open circuit potential of 1.222 V was determined. Data for these experiments appears in Fig. S4.†

### Conflicts of interest

The authors declare no competing financial interest.

### Acknowledgements

This research was supported as part of the Center for Molecular Electrocatalysis, an Energy Frontier Research Center funded by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences. Pacific Northwest National Laboratory is operated by Battelle for the U.S. Department of Energy.

### Notes and references

- 1 W. Lubitz, H. Ogata, O. Rüdiger and E. Reijerse, *Chem. Rev.*, 2014, **114**, 4081–4148.

2 Y. Nicolet and J. C. Fontecilla-Camps, *J. Biol. Chem.*, 2012, **287**, 13532–13540.

3 C. Madden, M. D. Vaughn, I. Díez-Pérez, K. A. Brown, P. W. King, D. Gust, A. L. Moore and T. A. Moore, *J. Am. Chem. Soc.*, 2012, **134**, 1577–1582.

4 D. L. DuBois, *Inorg. Chem.*, 2014, **53**, 3935–3960.

5 J. R. McKone, S. C. Marinescu, B. S. Brunschwig, J. R. Winkler and H. B. Gray, *Chem. Sci.*, 2014, **5**, 865–878.

6 J. R. McKone, N. S. Lewis and H. B. Gray, *Chem. Mater.*, 2013, **26**, 407–414.

7 V. S. Thoi, Y. Sun, J. R. Long and C. J. Chang, *Chem. Soc. Rev.*, 2013, **42**, 2388–2400.

8 P. Du and R. Eisenberg, *Energy Environ. Sci.*, 2012, **5**, 6012–6021.

9 W. J. Shaw, M. L. Helm and D. L. DuBois, *Biochim. Biophys. Acta, Bioenerg.*, 2013, **1827**, 1123–1139.

10 R. M. Bullock, A. M. Appel and M. L. Helm, *Chem. Commun.*, 2014, **50**, 3125–3143.

11 C. Tard and C. J. Pickett, *Chem. Rev.*, 2009, **109**, 2245–2274.

12 F. Gloaguen and T. B. Rauchfuss, *Chem. Soc. Rev.*, 2009, **38**, 100–108.

13 S. Ogo, K. Ichikawa, T. Kishima, T. Matsumoto, H. Nakai, K. Kusaka and T. Ohhara, *Science*, 2013, **339**, 682–684.

14 M. Y. Dahrenbourg and R. D. Bethel, *Nat. Chem.*, 2012, **4**, 11–13.

15 M. E. Carroll, B. E. Barton, T. B. Rauchfuss and P. J. Carroll, *J. Am. Chem. Soc.*, 2012, **134**, 18843–18852.

16 B. Ginovska-Pangovska, A. Dutta, M. L. Reback, J. C. Linehan and W. J. Shaw, *Acc. Chem. Res.*, 2014, **47**, 2621–2630.

17 N. Wang, M. Wang, J. Liu, K. Jin, L. Chen and L. Sun, *Inorg. Chem.*, 2009, **48**, 11551–11558.

18 J. M. Camara and T. B. Rauchfuss, *Nat. Chem.*, 2012, **4**, 26–30.

19 N. Wang, M. Wang, Y. Wang, D. Zheng, H. Han, M. S. G. Ahlquist and L. Sun, *J. Am. Chem. Soc.*, 2013, **135**, 13688–13691.

20 T. Liu, D. L. DuBois and R. M. Bullock, *Nat. Chem.*, 2013, **5**, 228–233.

21 J. M. Darmon, S. Raugei, T. Liu, E. B. Hulley, C. J. Weiss, R. M. Bullock and M. L. Helm, *ACS Catal.*, 2014, **4**, 1246–1260.

22 T. Liu, X. Wang, C. Hoffmann, D. L. DuBois and R. M. Bullock, *Angew. Chem., Int. Ed.*, 2014, **53**, 5300–5304.

23 K. T. Leffek, P. Pruszynski and K. Thanapalaingam, *Can. J. Chem.*, 1989, **67**, 590–595.

24 J. F. Coetzee and G. R. Padmanabhan, *J. Am. Chem. Soc.*, 1965, **87**, 5005–5010.

25 R. S. Nicholson and I. Shain, *Anal. Chem.*, 1964, **36**, 706–723.

26 J. M. Savéant, *Acc. Chem. Res.*, 1980, **13**, 323–329.

27 A. J. Bard and L. R. Faulkner, *Electrochemical Methods: Fundamentals and Applications*, Wiley, 2nd edn, 2000.

28 J. A. S. Roberts and R. M. Bullock, *Inorg. Chem.*, 2013, **52**, 3823–3835.

29 A. M. Appel and M. L. Helm, *ACS Catal.*, 2014, **4**, 630–633.

30 R. Custelcean and J. E. Jackson, *Chem. Rev.*, 2001, **101**, 1963–1980.

31 Z. M. Heiden, S. Chen, M. T. Mock, W. G. Dougherty, W. S. Kassel, R. J. Rousseau and R. M. Bullock, *Inorg. Chem.*, 2013, **52**, 4026–4039.

32 Bruker, *SAINT Program for Data Reduction*, Bruker AXS Inc., Madison, WI, USA, 2007.

33 Bruker, *SADABS Program for Absorption Correction of Area Detector Frames*, Bruker AXS Inc., Madison, WI, USA, 2007.

34 G. Sheldrick, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2008, **64**, 112–122.

35 O. V. Dolomanov, L. J. Bourhis, R. J. Gildea, J. A. K. Howard and H. Puschmann, *J. Appl. Crystallogr.*, 2009, **42**, 339–341.

36 J. P. Perdew, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1986, **33**, 8822–8824.

37 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, **78**, 1396.

38 D. Andrae, U. Häußermann, M. Dolg, H. Stoll and H. Preuß, *Theor. Chim. Acta*, 1990, **77**, 123–141.

39 A. Schäfer, H. Horn and R. Ahlrichs, *J. Chem. Phys.*, 1992, **97**, 2571–2577.

40 S. Chen, S. Raugei, R. Rousseau, M. Dupuis and R. M. Bullock, *J. Phys. Chem. A*, 2010, **114**, 12716–12724.

41 N. Kumar, D. M. Camaioni, M. Dupuis, S. Raugei and A. M. Appel, *Dalton Trans.*, 2014, **43**, 11803–11806.

42 S. Raugei, D. L. DuBois, R. Rousseau, S. Chen, M.-H. Ho, R. M. Bullock and M. Dupuis, *Acc. Chem. Res.*, 2015, **48**, 248–255.

43 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Chem. Theory Comput.*, 2009, **5**, 2447–2464.

44 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng, J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, Ö. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision B01*, Gaussian, Inc, Wallingford, CT, 2009.

45 P. B. Kisanga, J. G. Verkade and R. Schwesinger, *J. Org. Chem.*, 2000, **65**, 5431–5432.

