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# Electrocatalytic reduction of protons to dihydrogen by the cobalt tetraazamacrocyclic complex [Co(N<sub>4</sub>H)Cl<sub>2</sub>]<sup>+</sup>: mechanism and benchmarking of performances†‡

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The cobalt tetraazamacrocyclic  $[Co(N_4H)Cl_2]^+$  complex is becoming a popular and versatile catalyst for the electrocatalytic evolution of hydrogen, because of its stability and superior activity in aqueous conditions. We present here a benchmarking of its performances based on the thorough analysis of cyclic voltammograms recorded under various catalytic regimes in non-aqueous conditions allowing control of the proton concentration. This allowed a detailed mechanism to be proposed with quantitative determination of the rate-constants for the various protonation steps, as well as identification of the amine function of the tetraazamacrocyclic ligand to act as a proton relay during  $H_2$  evolution.

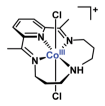
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Molecular cobalt complexes are popular and versatile catalysts for the electrocatalytic evolution of hydrogen.1-5 Recently, the cobalt complex [Co(N<sub>4</sub>H)Cl<sub>2</sub>]<sup>+</sup> (Cat1, Fig. 1) based on the tetraazamacrocyclic 2,12-dimethyl-3,7,11,17-tetraazabicyclo [11.3.1]heptadeca-1(17),2,11,13,15-pentaene ligand,6 described by Karn and Busch in 1966, has received increased interest<sup>7-9</sup> namely because this catalyst proves active and stable for the evolution of H<sub>2</sub> from fully aqueous solutions.<sup>7-17</sup> A study carried out under homogeneous conditions using chemical reductants or photochemical activation confirmed the superior activity of Cat1 in fully aqueous media,18 and X-ray absorption spectroscopic monitoring of a homogeneous photocatalytic system for  $H_2$  evolution based on Cat1 indicated an ECEC mechanism (E = monoelectronic electrochemical reduction, C = protonation step) starting from the bisaqua Co(II) complex. 13,14 However, very few metrics are currently available to benchmark the catalytic activity of this compound. **Cat1** was included in a benchmarking study for  $H_2$ -evolving electrocatalysts carried out in aqueous electrolyte<sup>19</sup> but under quite acidic conditions likely to induce reductive degradation of the ligand during the test and formation of metallic particles responsible for the observed HER activity.<sup>20,21</sup> To gain quantitative information on the  $H_2$  evolution mechanism mediated by **Cat1**, we therefore revisited the non-aqueous conditions investigated by Lau and coworkers,<sup>22</sup> where it is easier to control the concentration and chemical potential of protons.<sup>23</sup>

The cyclic voltammogram of Cat1 (perchlorate salt) in CH<sub>3</sub>CN (with 0.1 M  $^n$ Bu<sub>4</sub>NBF<sub>4</sub>, Fig. 2 and S1‡) displays two quasi-reversible systems at -0.47 V ( $\Delta E_p = 92$  mV) and -0.96 V ( $\Delta E_p = 86$  mV)  $\nu s$ . Fc<sup>+</sup>/Fc, corresponding to the Co<sup>III/II</sup> and Co<sup>III/I</sup> redox processes, respectively, based on previous literature. Of note, we will formally use the Co<sup>II</sup> notation throughout this article, while the electronic state could also correspond to a reduced N<sub>4</sub>H ligand ( $\pi$ -radical anion) antiferromagnetically



 $[Co(N_4H)Cl_2]^+$  (Cat1)

Fig. 1 Structure of Cat1.

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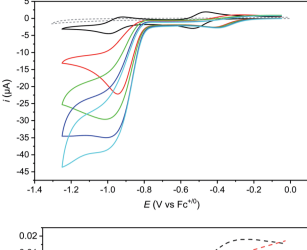
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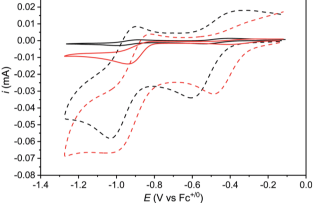


Fig. 2 Top: cyclic voltammograms of Cat1 (0.5 mM) in CH $_3$ CN (+0.1 M  $^{\prime\prime}$ Bu $_4$ NBF $_4$ ) recorded at a glassy carbon electrode (1.6 mm diameter) in the absence (black trace) and in the presence of 5.0 (red trace), 10.0 (green trace), 12.5 (navy trace) and 15.0 mM (cyan trace) p-cyanoanilinium tetrafluoroborate; scan rate: 100 mV s $^{-1}$ . A control voltammogram of 15 mM p-cyanoanilinium tetrafluoroborate without Cat1 (gray dashed trace) is shown for comparison; bottom: cyclic voltammograms of Cat1 (0.5 mM) in CH $_3$ CN (+0.1 M  $^n$ Bu $_4$ NBF $_4$ ) recorded at a glassy carbon electrode in the absence (black) and in the presence (red) of 5 mM p-cyanoanilinium tetrafluoroborate; scan rate: 0.1 (solid) and 10 V s $^{-1}$  (dashed). See Fig. S5 $\ddagger$  for control voltammograms without catalyst under similar conditions.

coupled to a low-spin Co(II) ion.<sup>24</sup> A third ligand-centered process is observed at -1.89 V ( $\Delta E_{\rm p}=69$  mV)  $\nu s$ .  ${\rm Fc}^+/{\rm Fc}^{+2}$  Upon addition of p-cyanoanilinium tetrafluoroborate ( $pK_{\rm a}=7.0$  in  ${\rm CH_3CN})^{25}$  acting as a proton source, a catalytic wave develops on the top of the  ${\rm Co^{II/I}}$  wave as previously described.<sup>22</sup> This electrocatalytic behavior corresponds to  ${\rm H_2}$  evolution with >90% faradaic yield<sup>22</sup> and nicely mirrors the one observed for  ${\rm Cat1}$  in mildly acidic aqueous solution.<sup>11,13,22</sup> The addition of stronger acids such as  ${\rm HBF_4 \cdot Et_2O}$  or  ${\rm CF_3SO_3H}$  also triggers  ${\rm H_2}$  evolution catalysis but is detrimental to the stability of  ${\rm Cat1}$  at high concentration.  ${\rm Cat1}$  is unable to catalyze the reduction of acids with higher  $pK_a$  values, starting with p-toluenesulfonic acid ( $pK_a=8.3$  in  ${\rm CH_3CN}$ ).<sup>26</sup>

Noteworthily, the addition of acid also affects the Co<sup>III/II</sup> system, which shifts to more positive potentials and partly loses reversibility. <sup>1</sup>H NMR experiments confirmed that the Co<sup>III</sup> form of Cat1 is not protonated under these conditions (Fig. S2‡),

neither does p-cyanoaniline coordinate to any of the Co<sup>III</sup>, Co<sup>II</sup> and Co<sup>I</sup> states of Cat1 (Fig. S3‡) in the absence of acid. However, when 5 equivalents of p-cyanoanilinium tetrafluoroborate is added to an electrochemically-generated solution of the CoII form of Cat1, the EPR signal is significantly changed, from a broad signal centered at  $g_1 = 2.240$ ,  $g_2 = 2.130$  and  $g_3 = 2.004$ to a better resolved spectrum characteristic for a low spin d<sup>7</sup> (S  $=\frac{1}{2}$ ) electronic configuration of Co (Fig. 3). It furthermore displays a 5-line superhyperfine structure with an intensity ratio of 1:2:3:2:1 and a coupling constant of 45 Hz in line with the coordination of two equivalent nitrogen-based ligands. Of note, a similar behavior was observed when the CoII form of Cat1 was prepared by chemical reduction with cobaltocene instead of exhaustive bulk electrolysis (Fig. S4‡). No such change is observed when p-cyanoaniline is added (Fig. S4<sup>‡</sup>). When HBF<sub>4</sub>·Et<sub>2</sub>O is used as proton source instead of p-cyanoanilinium tetrafluoroborate, this superhyperfine structure is changed to a 3-line structure with a coupling constant of 40 Hz (Fig. 3), suggesting the coordination of a single nitrogen-based axial ligand, CH<sub>3</sub>CN being the only plausible one under these conditions. In the former case, coordination of CH3CN and pcyanoanilinium (or p-cyanoaniline generated in situ upon protonation of the complex) can be considered without being possible to discriminate one from the other at the EPR level. Taken all together, these observations suggest (i) protonation of the ligand in Cat1 occurs upon reduction from the Co<sup>III</sup> to the Co<sup>II</sup> state in the presence of acid; (ii) at the same time, chloride axial ligands are displaced for nitrogen ligands; (iii) the nature and number of axial ligands depend on the nature of the acid employed. It should also be underlined that the binding of one vs. two ligands in the cobalt + II oxidation state likely depends on a subtle balance of their donating ability, as previously stated

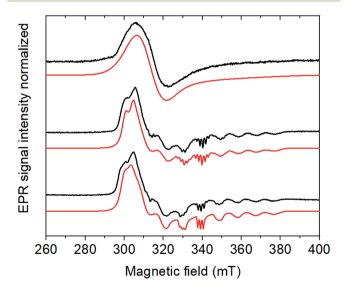


Fig. 3 CW X-band EPR spectra (9.65 GHz) of the electrochemically generated  $\mathrm{Co}^{II}$  form of Cat1 (0.5 mM) as prepared (top), with 5 eq. of the acids p-cyanoanilinium tetrafluoroborate (middle), HBF<sub>4</sub> (bottom) and their respective simulations in red. Experimental conditions: 30 K, 1 mW microwave power, 1600 G field sweep. Simulation parameters are reported in Table S1.‡

for Cat1 (ref. 11) and also found for cobaloximes and related cobalt diimine-dioxime complexes.27-30 A possible structure of Co<sup>II</sup>LH, in line with a previous proposition, <sup>13</sup> is shown in Scheme 1.

Although the formation of the protonated Co<sup>II</sup> derivative of Cat1 falls into the proton-coupled electron transfer classification, the cathodic peak potential does not follow the increase of 29 mV per decade of acid concentration expected for an irreversible EC process with a fully displaced protonation equilibrium (e.g. in the pure kinetic KP zone);31 rather the cathodic peak potential rapidly shifts to a new value upon addition of acid and then keeps this value unchanged upon further addition (Fig. 2). This behavior is characteristic of the extraordinary kinetic (KE) zone,31 with fast protonation of the CoII species so that, even with few equivalents of acid added, the new wave is observed at a potential close to the apparent standard potential of the  $Co^{III}/Co^{II}LH$  couple, thus with a  $\sim$ 250 mV shift compared to the original Co<sup>III</sup>/Co<sup>II</sup> couple. Recording the cyclic voltammograms at a significantly higher scan rate (10 V s<sup>-1</sup>) did not allow approaching the pure kinetic KP zone although a slightly more progressive evolution of the cathodic peak potential was observed with an increase of ~200 mV per decade of acid

Cat1 CollHLH Co<sup>III</sup>HLH

Scheme 1 Proposed ECEC mechanism for H<sub>2</sub> evolution mediated by Cat1. L and L' indicate acetonitrile or p-cyanoaniline.

concentration (Table S2‡). Simulations using the DigiElch software allowed to reproduce the potential shift of such an EC process at both 0.1 and 10 mV s<sup>-1</sup> using a protonation equilibrium constant higher than 104 and a bimolecular protonation rate of  $10^7$  mol<sup>-1</sup> L s<sup>-1</sup>, suggesting that the amine moiety can potentially act as a proton relay during catalysis.32

Protonation of the CoII state should also alter the standard potential of the Co<sup>II</sup>/Co<sup>I</sup> couple. At 100 mV s<sup>-1</sup>, this redox process is hidden by the catalytic wave it triggers. However, measuring cyclic voltammograms at 10 V s<sup>-1</sup> enabled the catalysis to be outrun and a reversible wave to be recovered centred at -0.89 V vs. Fc+/Fc (Fig. 2 bottom) that likely corresponds to the actual CoILH/CoILH couple responsible for catalysis. Remarkably, raising acid concentration, we found that catalysis proceeds under pure kinetic conditions, where a catalytic current plateau, independent of the scan rate, is observed (Fig. 4) because substrate consumption is negligible (pure kinetic KS zone).31,33 Under these conditions, the mid-wave potential of the catalytic process is found at -0.87 V vs. Fc<sup>+</sup>/ Fc. Varying the catalyst concentration for a given acid concentration showed a linear dependence of the catalytic current with the catalyst concentration (Fig. S6‡). The catalytic plateau current also linearly varies with the acid concentration (Fig. S7<sup>‡</sup>).

Together these data are consistent with a mechanism for H<sub>2</sub> evolution catalyzed by Cat1 following an ECEC reaction scheme (Scheme 1), where E and C stand for electrochemical steps and chemical (i.e. protonation) steps and with the second reduction occurring at a potential more positive to that of the first one: reduction of the protonated Co<sup>II</sup> (Co<sup>II</sup>LH) state yields Co<sup>I</sup>LH, which is further protonated to yield the Co<sup>III</sup>HLH species. Further reduction of this derivative then produces the CoII hydride Co<sup>II</sup>HLH species that is further protonated to evolve H<sub>2</sub> and regenerate the starting Co<sup>II</sup>LH complex. Some of the

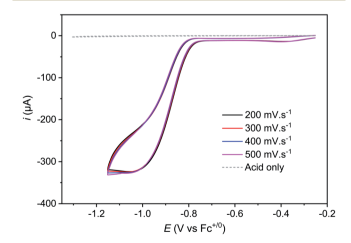


Fig. 4 Cyclic voltammograms of Cat1 (1.6 mM) in CH<sub>3</sub>CN (+0.1 M <sup>n</sup>Bu<sub>4</sub>NBF<sub>4</sub>) recorded at a glassy carbon electrode (1.6 mm diameter) in the presence of 60 mM p-cyanoanilinium tetrafluoroborate at various scan rates, scan rate ranging from 200 to 500 mV  $\rm s^{-1}$  and with ohmic drop compensation. A control voltammogram of 60 mM p-cyanoanilinium tetrafluoroborate without Cat1 at 100 mV  $s^{-1}$  is shown for comparison.

distorded geometries displayed in Scheme 1 for transient intermediates involved in the catalytic cycle are approximate and we do realize that they do not correspond to ideal geometries for coordination complexes with corresponding electronic structures. Protonation of the amine residue of the macrocyclic ligand namely generates an ammonium moiety unable to coordinate the cobalt center, though with significant steric hindrance preventing for example the adoption of a perfect square planar (or octahedral) geometry favored by the d<sup>8</sup> (or d<sup>6</sup>) configuration in the CoILH (or CoIIIHLH) species, as demonstrated for the unprotonated Co<sup>I</sup> (ref. 14) or Co<sup>III</sup>-hydride species,34 respectively. Of note this family of cobalt complexes can accommodate various coordination spheres as demonstrated by the heptacoordinated systems recently reported.35,36

Eqn (1) gives the plateau current  $i_p$  for such a process, assuming that the two electrons required to complete catalytic turnover are transferred from the electrode to the catalyst.

$$i_{\rm pl} = 2FSC_{\rm cat}^0 \sqrt{k_2 D_{\rm cat} C_{\rm AH}^0} \tag{1}$$

where F is the Faraday constant, S is the geometric electrode surface area,  $C_{\text{cat}}^0$  is the concentration of the catalyst and  $D_{\text{cat}}$  is the diffusion coefficient of the catalyst, determined to be 10<sup>-5</sup> cm<sup>2</sup> s<sup>-1</sup> from the scan-rate dependence of the peak current of the first cathodic wave and the Randles-Sevcik equation (see ESI‡).

From plateau currents measured over a range of concentrations for both catalyst and acid (Fig. S6 and S7‡), we found a value of 5.3  $\pm$  0.1  $\times$  10<sup>3</sup> mol<sup>-1</sup> L s<sup>-1</sup> for the second order rate constant  $k_2$ . The rate constant for the first protonation step  $k_1$  of ECEC processes is accessible from the Foot-of-the-Wave Analysis (FOWA).37,38 This analysis requires the knowledge of the apparent standard potential of the redox couple involved, i.e.  $Co^{II}LH/Co^{I}LH$ , which could be determined to  $-0.89 \text{ V} \text{ vs. } Fc^{+}/Fc$ using high scan rates (Fig. 2 bottom). FOWA was performed at 3 different scan rates (100, 400 and 1000 mV s<sup>-1</sup>) for two different acid concentrations (5 and 25 mM) and gave a value of 2.5  $\pm$  0.4  $\times$  10<sup>4</sup> mol<sup>-1</sup> L s<sup>-1</sup> for the second order rate constant  $k_1$ (Fig. S8‡). The higher value of  $k_1$  compared to  $k_2$  is in line with the midwave of the catalytic process being shifted positively compared to the standard potential of the Co<sup>II</sup>LH/Co<sup>I</sup>LH couple.<sup>38</sup> Analysis of this shift using eqn (2) for the data shown in Fig. S6 and S7‡ also leads to an average value of  $2.5 \times 10^4$  $\text{mol}^{-1} \text{L s}^{-1}$  for the second order rate constant  $k_1$ , although with a much larger error margin.

$$E_{\text{cat/2}} = E_{\text{Co}^{\text{II}}\text{LH}/\text{Co}^{\text{I}}\text{LH}}^{0} + \frac{RT}{2F} \ln \frac{k_{1}}{k_{2}}$$
 (2)

The  $k_1$  value can finally be confirmed from the analysis of cyclic voltammograms recorded at low acid concentration where the system belongs to the "total catalysis" regime (Fig. 5).31,33 In this regime, catalysis is so fast that all the acid present in the diffusion layer is consumed during the sweep of the catalytic wave. As a consequence, the unprotonated Co(II) form of Cat1 is regenerated after catalysis and its reduction is observed at -0.96 V vs. Fc<sup>+</sup>/Fc.

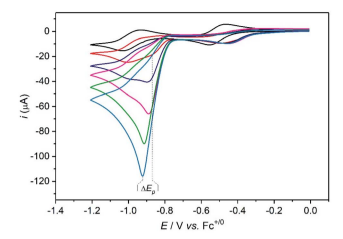


Fig. 5 Cyclic voltammograms of Cat1 (2 mM) in CH<sub>3</sub>CN (+0.1 M  $^{n}$ Bu<sub>4</sub>NBF<sub>4</sub>) recorded at a glassy carbon electrode (1.6 mm diameter) in the presence of various concentrations of p-cyanoanilinium tetrafluoroborate at 100 mV s $^{-1}$ : 0 (black), 4 (red), 8 (navy), 12 (magenta), 16 (green), 20 (cyan) mM.

Based on simulations, Dempsey and coworkers could propose eqn (3) to describe the variation of the catalytic peak potential  $E_{\rm p}$ .<sup>39</sup>

$$E_{\rm p} = E_{\rm p}^0 - 0.409 \frac{RT}{F} + \frac{RT}{2F} \ln \left( \frac{RT}{F_{\rm v}} \frac{D_{\rm cat}}{D_{\rm AH}} \frac{k_1 (2C_{\rm cat}^0)^2}{C_{\rm AH}^0} \right)$$
(3)

As the intensity of the catalytic wave in the "total catalysis" regime is controlled by the diffusion of the acid, the  $D_{cat}/D_{AH}$ ratio between the diffusion coefficient of the catalyst and pcyanoanilinium cation can be obtained from the ratio between the catalytic peak current and the current of the monoelectronic Co<sup>II</sup>/Co<sup>I</sup> wave measured in the absence of acid, according to eqn (4).

$$\frac{i_{\text{peak}}}{i_{\text{p}}^{0}} = 1.365 \frac{C_{\text{AH}}^{0}}{C_{\text{cat}}^{0}} \sqrt{\frac{D_{\text{AH}}}{D_{\text{cat}}}}$$
 (4)

Applying eqn (3) to the data shown in Fig. 5 yields a  $k_1$  value of 2.05  $\pm$  0.30  $\times$  10<sup>4</sup> mol<sup>-1</sup> L s<sup>-1</sup> for the second order rate constant  $k_1$ . This value is in good agreement with the one obtained by FOWA although neither  $k_1$  nor  $k_2$  comply with the condition of being greater than  $10^7 \, \text{mol}^{-1} \, \text{L s}^{-1}$  required for eqn (3) to be valid.<sup>39</sup>

Interestingly, these data allow ruling out homolytic H2 evolution mediated by Cat1. In such a mechanism, the first protonation step generating the  $Co^{III}$ -H derivative with the  $k_1$ rate constant still exists but it is followed by reductive elimination of  $H_2$  from two  $Co^{III}$ -H species with a  $k_d$  second-order rate constant. The linear dependency of the plateau current with the catalyst concentration (Fig. S6‡) rules out homolytic H<sub>2</sub> evolution from this CoIII-H derivative in the non-steady state where the rate-determining step is the reductive elimination step. 38,40 The identification of two distinct rate constants with  $k_1$ being the largest one also allows to rule out steady state

homolytic H<sub>2</sub> evolution with the rate-determining step being the formation of the Co<sup>III</sup>-H derivative, therefore implying that  $k_1$  <  $k_{\rm d}$ . Finally, while the mechanism proposed in Scheme 1 does not formally exclude that formation of the Co<sup>III</sup>HLH species may proceed through intramolecular protonation of the Co<sup>I</sup> center of Co<sup>I</sup>LH followed by reprotonation of the ligand, we believe that such a possibility is unlikely as Et<sub>3</sub>NH<sup>+</sup>, an acid with a p $K_a$  similar to that of the protonated  $N_4H_2^+$  ligand in  $Co^{I}LH$ , is unable to protonate  $Co^{I}$  complexes with similar  $E^{0}_{Co^{II}/Co^{I}}$  value such as [Co<sup>I</sup>(dmgBF<sub>2</sub>)<sub>2</sub>(CH<sub>3</sub>CN)] (dmgH<sub>2</sub> = dimethylglyoxime) in CH<sub>3</sub>CN.41

The mechanism proposed here for H<sub>2</sub> evolution differs from the one proposed by Llobet and Gimbert-Suriñach for aqueous conditions.13,14 First we clearly evidenced that the CoII form of Cat1 is protonated under the conditions investigated here. Ligand protonation also occurs in aqueous electrolyte, as shown by a 59 mV pH unit <sup>-1</sup> shift of the redox process reported by Peters and coworkers. 19 Second, we propose that H<sub>2</sub> is formed from a CoII-H and not a CoIII-H intermediate. We recognize that the electrochemical responses of ECEC and ECCE sequences, both starting from a CoII derivative and implying Co<sup>II</sup>-H and Co<sup>III</sup>-H active species, respectively, are similar. However DFT calculations clearly demonstrated that the standard potential of the Co<sup>III</sup>-H/Co<sup>II</sup>-H is more positive than that of the Co<sup>II</sup>/Co<sup>I</sup> couple, <sup>13</sup> a feature also observed for cobaloximes and cobalt diimine-dioxime complexes for which the ECEC mechanism is now accepted. Importantly, the mechanism shown in Scheme 1 involves the Co<sup>II</sup>HLH species that was proposed by Llobet and Gimbert-Suriñach under photocatalytic conditions,13 therefore unifying the mechanistic understanding of this catalyst. DFT calculations previously indicated a nearthermoneutral intramolecular H2 evolution step from this protonated hydride intermediate.13 Our analysis shows that this step is also the rate-determining one, which explains why catalysis is so fast. Still, the observation that the rate constant of this step is first order in acid concentration suggests that intramolecular H2 formation is coupled with protonation, either in a concerted manner or through kinetic coupling with the fast reprotonation of the amine group of the ligand.

The maximal turnover frequency (TOF<sub>max</sub>) of Cat1 therefore approximates  $k_2 \times [\text{acid}]$  and a TOF<sub>max</sub> value of 5.3  $\times$  10<sup>3</sup> s<sup>-1</sup> can be extrapolated for 1 M p-cyanoanilinium tetrafluoroborate concentration. Based on this value and an apparent equilibrium potential of the  $H^+/H_2$  couple of -0.47 V vs.  $Fc^+/Fc$  at 1 M pcyanoanilinium tetrafluoroborate concentration and taking homoconjugation into account,26 we could derive the red trace in the catalytic Tafel plot42,43 shown in Fig. 6. Interestingly, Cat1 displays significant catalytic activity  $(\log(TOF/s^{-1}) > 1)$  at low overpotential values, a property shared by very few other molecular complexes including cobaloximes, 42,44 DuBois' nickel bisdiphosphine catalysts<sup>42,45</sup> and bis(thiosemicarbazone) cobalt46 and nickel47 complexes. Its overpotential requirement, estimated to be  $\sim$ 400 mV at the catalytic half-wave potential (and corresponding to the inflexion point in the red catalytic Tafel plot on Fig. 6) is  $\sim$ 100 mV higher than that of cobaloxime, cobalt diimine-dioxime48 or DuBois' complexes. Importantly, the high TOFmax value also places Cat1 in an intermediate

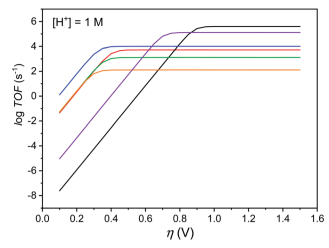


Fig. 6 Catalytic Tafel plots. Comparison of performances for H<sub>2</sub> evolution catalyzed by Cat1 in CH3CN in the presence of 1 M p-cyanoanilinium (red line), with other catalysts reported in the literature: black: Fe<sup>II</sup>TPP, DMF, Et<sub>3</sub>NH<sup>+</sup>;<sup>42,49</sup> blue: Co<sup>II</sup>(dmgH)<sub>2</sub>py, DMF, Et<sub>3</sub>NH<sup>+</sup>;<sup>42</sup> green:  $[Ni^{II}(P_2^{Ph}N_2^{C_6H_4X})_2]^{2+}$ ,  $X = CH_2P(O)(OEt)_2$ , MeCN, DMFH<sup>+</sup>;<sup>42,45</sup> orange: 4-{bis[4-(p-methoxyphenyl)thiosemicarbazone]}-2,3-butane cobalt, DMF, Et<sub>3</sub>NH<sup>+</sup>;<sup>46</sup> purple: [Co<sup>II</sup>(bapbpy)Cl], DMF, Et<sub>3</sub>NH<sup>+</sup>.<sup>23</sup>

position between cobaloximes and DuBois' complexes, the two champion H2-evolving molecular catalyst series identified so far in non-aqueous solvents.42 It is noteworthy that all three catalysts possess proton relays in their second-coordination sphere and the protonation of these relays is coupled with the metalcentered reduction step that sets the potential of the catalytic wave. The same conclusion can be reached with thiosemicarbazone nickel and cobalt complexes, with ligandcentered reduction occurring in these cases at quite positive potentials.46,47

#### Conclusions

While Cat1 is becoming more and more popular as a molecular H<sub>2</sub>-evolving catalyst for the design of aqueous systems, it is increasingly important to advance the understanding of its H<sub>2</sub> evolution catalysis mechanism13 and performance by providing insight into the catalytic steps involved. This is especially the case when considering structural modification50 or molecular engineering<sup>51</sup> in order to either enhance catalytic activity or stability16 or immobilise a catalytic centre for integration in photoelectrodes<sup>7-9</sup> or devices. In the societal context, both of these objectives are ultimately necessary to achieve industrial relevancy and technological maturity of hydrogen-producing electrolysers based on molecular catalysts made from earthabundant elements.2,4

In this study, new EPR evidence for the structure of the Co<sup>II</sup> state of the catalyst and its dependence on the presence and nature of the acid has been presented, as well as NMR and cyclic voltammetry data indicating that fast protonation of the ligand occurs at the Co<sup>II</sup> stage. Carrying on from previous work in the literature, 13,14,22 these results reconcile mechanisms at play under electrochemical and photochemical conditions.

Furthermore, a proper utilisation of the most advanced analytical methods available for molecular catalysis of electrochemical reactions has allowed definitively ruling out a homolytic H<sub>2</sub> evolution mechanism and substantiated the proposed heterolytic ECEC mechanism, for which the rate constants for the two successive protonation steps could be determined. The rate-determining step has been confirmed as the intramolecular H<sub>2</sub> evolution step, surmised to be coupled to the second protonation, thereby regenerating a protonated ligand and thus acting as a proton relay in catalysis,<sup>32</sup> which, interestingly, is at variance with the behaviour of the dioxime bridge in cobalt diimine–dioxime complexes.<sup>48</sup> Catalytic Tafel plots could be derived to enable the benchmarking of the H<sub>2</sub> evolution performance of **Cat1** alongside other highly efficient catalysts and confirm its place on the podium.

#### Conflicts of interest

There are no conflicts to declare.

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