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Tetraphenylporphyrin electrocatalysts for the hydrogen evolution reaction: applicability of molecular volcano plots to experimental operating conditions†

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Recent years have seen an increasing interest in molecular electrocatalysts for the hydrogen evolution reaction (HER). Efficient hydrogen evolution would play an important role in a sustainable fuel economy, and molecular systems could serve as highly specific and tunable alternatives to traditional noble metal surface catalysts. However, molecular catalysts are currently mostly used in homogeneous setups, where quantitative evaluation of catalytic activity is non-standardized and cumbersome, in particular for multi-step, multielectron processes. The molecular design community would therefore be well served by a straightforward model for prediction and comparison of the efficiency of molecular catalysts. Recent developments in this area include attempts at applying the Sabatier principle and the volcano plot concept – popular tools for comparing metal surface catalysts – to molecular catalysis. In this work, we evaluate the predictive power of these tools in the context of experimental operating conditions, by applying them to a series of tetraphenylporphyrins employed as molecular electrocatalysts of the HER. We show that the binding energy of H and the redox chemistry of the porphyrins depend solely on the electron withdrawing ability of the central metal ion, and that the thermodynamics of the catalytic cycle follow a simple linear free energy relation. We also find that the catalytic efficiency of the porphyrins is almost exclusively determined by reaction kinetics and therefore cannot be explained by thermodynamics alone. We conclude that the Sabatier principle, linear free energy relations and molecular volcano plots are insufficient tools for predicting and comparing activity of molecular catalysts, and that experimentally useful information of catalytic performance can still only be obtained through detailed knowledge of the catalytic pathway for each individual system.

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1 Introduction

Production of H₂ fuel through water electrolysis is one of many important steps towards a more sustainable society. This process requires robust redox catalysts to be economically and environmentally feasible. Pt has long been known as the state-of-the art catalyst of the hydrogen evolution reac-

tion (HER), but as it is also scarce and expensive,^{1–3} much attention is turning towards homogeneous, molecular alternatives. Molecular catalysts can be produced from more earth-abundant elements,⁴ have well-defined catalytic sites^{5,6} and can be tuned towards high activity and selectivity through molecular design.^{4,5} Several molecular systems, such as water-soluble transition metal complexes,⁷ Ni(II) bi(diphosphine) complexes,⁸ and various porphyrins,^{9–14} have already been shown to be catalytically active towards the HER. However, molecular systems can currently rarely compete with heterogeneous, industrial catalysts in terms of stability and lifetime. In addition, complex reaction mechanisms and a lack of consensus on how to best evaluate reaction kinetics mean that development of improved molecular catalysts often requires cumbersome testing of individual designs. The study of molecular catalysis would therefore greatly benefit from a straightforward tool to predict and compare catalytic properties.

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In heterogeneous catalysis, the Sabatier principle is a recognized model for comparing and selecting catalytic materials.¹⁵ It states that if the substrate binds too weakly to the catalyst, there will be an energy barrier between the initial state and the following intermediate, and if the substrate binds too strongly, there will be an energy barrier between the intermediate and the next step or product. The Sabatier principle has been used to confirm and predict relative efficiencies of metal surfaces used for catalysis of the hydrogen evolution,^{16–18} oxygen reduction^{19,20} and oxygen evolution reactions.^{21,22} By plotting a measure of the catalytic activity against the binding energy of a particular intermediate, a so-called volcano plot can be produced, where the most efficient catalyst is located at the apex.

According to the Sabatier principle, the free-energy landscape of an ideal catalyst is flat at equilibrium, meaning that at equilibrium there are no energy barriers between the substrate, the reaction intermediates and the product. For hydrogen evolution on metal surfaces, construction of a volcano plot representing a flat energy landscape is straightforward, given that hydrogen bound to the surface is considered the only intermediate.^{15,17,23} The binding energy of hydrogen, $\Delta\epsilon_{\text{H}}$, is then a good descriptor of catalytic efficiency.^{17,18} For catalytic cycles involving more steps, the free energy change of one step is not necessarily independent of the free energy change of the others. For example, the binding energies of OH and OOH in the oxygen evolution reaction have been shown to be linearly related.^{20–22} In fact, assuming or identifying linear free energy dependencies between different intermediates gives rise to so-called scaling relations, which can be used to produce volcano plots and optimize energy barriers for a series of catalysts.^{15,20–22,24}

In molecular homogeneous catalysis, scaling relations are often used to illustrate the trade-off between catalytic activity and overpotential.^{25–27} Recently, such relations have been used to produce molecular volcano plots.^{25,28,29} However, these plots have been realized for enzymes³⁰ or in computational settings, and have not yet been fully connected to experimental studies of small molecular catalysts. In addition, the Sabatier analysis and volcano plots in both heterogeneous and homogeneous setups tend to focus on the thermodynamic aspects of catalysis. Indeed, free energy relations and so-called theoretical minimum overpotentials reveal nothing of the inevitable kinetic aspects of real-world catalysis, and it has been suggested on numerous occasions that thermodynamic profiles and/or linear free energy relations are not sufficient for accurate prediction of catalytic quality.^{26,31–38}

In this work, we investigate the value of applying the Sabatier principle, linear scaling relations and the volcano plot concept to molecular catalysis in relation to experimental reaction kinetics. We report the electrochemistry and catalytic activity towards the HER for a series of tetraphenylporphyrins, and evaluate the applicability of the Sabatier principle by comparing the computationally derived $\Delta\epsilon_{\text{H}}$ to various experimental electrochemical kinetic parameters. We show that $\Delta\epsilon_{\text{H}}$ abides by a linear scaling relation between the initial steps of the catalytic reaction, and that it agrees with the thermo-

dynamic profiles of the series. We also show that the binding energy is not sufficient to accurately predict reaction kinetics and catalytic activity. We conclude that while the Sabatier principle and the volcano plot concept may be transferrable to molecular catalysis in terms of pure thermodynamics, they cannot be used to determine which catalyst will perform best in a real-world setting. For successful application of molecular design, other tools must be developed.

2 Experimental section

2.1 General

The series of molecular catalysts used in this study consists of free base and metalated *meso*-tetraphenylporphyrins, referred to as MTPP. Unless otherwise stated, the metals are in oxidation state +II. NiTPP, CuTPP, CoTPP and ZnTPP were purchased from PorphyrChem and used without further purification. Al(III)TPPCL, FeTPP, MgTPP and O=Ti(IV)TPP were synthesized through metalation of commercially available H₂TPP as previously reported³⁹ and AgTPP is also commercially available.

Dichloromethane (DCM, puriss. p.a., ACS reagent, reagent grade, ISO, GC, ≥99.9%, VWR chemicals) was used as the solvent for all electrochemical and spectroscopic studies. Tetrabutylammonium hexafluorophosphate (TBAPF₆), purchased from Sigma-Aldrich, was used as received as the supporting electrolyte. Glassy carbon electrodes, reference electrodes and Pt wire electrodes were purchased from Bio-Logic.

2.2 Electrochemical characterization

Each porphyrin was dissolved in 0.1 M TBAPF₆/DCM; H₂TPP and ZnTPP to 2.5 mM and all other porphyrins to saturation. The solution was deoxygenated by purging with N₂ gas. Cyclic and square wave voltammetry were then performed under N₂ atmosphere in a three-electrode setup using an Autolab PGSTAT032N potentiostat from Metrohm AG. A glassy carbon disc electrode (3 mm diameter, 70 mm long) was used as the working electrode, a Pt wire as the counter electrode and the Ag/Ag⁺ couple as the reference electrode. The reference electrode was kept in a separate compartment connected to the bulk solution by a porous frit, and the Fc/Fc⁺ couple was used as an internal standard. Unless otherwise stated, all cyclic voltammograms were performed at 0.1 V s^{−1}. Square wave voltammetry was performed at 15 Hz, using a 4 mV potential increment and an amplitude of 25 mV.

2.3 UV/vis spectroscopy

UV/vis spectroscopy was performed using an Agilent 8453 UV-visible single beam spectrophotometer and a 1 mm thin layer spectroelectrochemical quartz cell. Each porphyrin was dissolved to 25 μM in DCM and the absorbance spectrum recorded between 190 and 1100 nm. For each porphyrin, a blank spectrum was recorded against porphyrin free solution and subtracted from the sample spectrum. Absorbance spectra



were also recorded with 22 equivalents of benzoic acid (BzOH) added to both the blank and the sample spectrum.

2.4 Catalysis of hydrogen evolution

The catalytic activity of the porphyrins towards the HER was evaluated using the same three-electrode setup as for cyclic voltammetry, but using a porphyrin concentration of 0.2 mM in 0.1 M TBAPF₆/DCM. 0.1 M BzOH/DCM was used as the proton source, as BzOH is not strong enough to cause demetallation of the porphyrins.¹¹ The amount of acid was stepwise increased by adding 1 equivalent at a time to 20 mL porphyrin solution. Cyclic voltammograms (CVs) were recorded at 1–10 equivalents of acid, and linear sweep voltammograms (LSVs) at 1–22 equivalents of acid. Linear sweep voltammetry was also performed at 1–22 equivalents of acid in porphyrin-free solution to exclude catalytic effects of the glassy carbon electrode, and cyclic voltammetry was conducted before and after each round of catalytic experiments to ensure adsorbed species did not contribute to the catalytic process.

3 Computational details

All computations were performed using density functional theory (DFT) on the Tetralith cluster provided by the Swedish National Infrastructure for Computing (SNIC). Gaussview 6.1.1. was used to build the molecules, and their electronic structures were evaluated using the Becke, 3-parameter, Lee–Yang–Parr functional⁴⁰ with Grimme's empirical dispersion correction⁴¹ (B3LYP-D3) as implemented in Gaussian 16. The geometry of each compound was optimized in gas phase using the 6-31++(d,p) basis set^{42,43} for all elements lighter than and including Ar, and using Stuttgart/Dresden effective core potentials⁴⁴ on all elements heavier than Ar. Zero-point energies and thermal corrections to reaction free energies were obtained at 298.15 K and 1 atm through frequency calculations. To improve the accuracy of electronic and Gibbs free energies, single point calculations were performed using the 6-311++(d,p)^{42,45} basis set for elements lighter than and including Ar, and the def2-TZVP⁴⁶ basis set for elements heavier than Ar. Atomic charges according to Charge Model 5 (CM5),⁴⁷ as well as Mulliken spin densities, were obtained from the single point calculations. No solvation model was used, as the effects of solvation are expected to more or less cancel when calculating binding energies, meaning that solvation will have negligible effects on the trends presented here.

As transition metals can occur in several spin states, each structure except the free base, AlTPP and MgTPP was optimized at its three lowest multiplicities (singlet, triplet and quintet, or doublet, quadruplet and sextet). Reaction free energies and other inferred quantities were calculated using the multiplicity that resulted in the lowest energy including thermal corrections. Porphyrins that undergo metal reduction before ligand reduction, or metal oxidation before ligand oxidation, were optimized with a total molecular charge of –1 or +1, respectively.

4 Results and discussion

4.1 Electrochemical characterization

The porphyrins studied in this work are shown in Fig. 1. Electrochemical characterizations were obtained for AgTPP, Al(III)TPP, CoTPP, Co(III)TPP, CuTPP, FeTPP, H₂TPP, MgTPP, NiTPP, O=Ti(IV)TPP and ZnTPP. Catalytic properties were evaluated experimentally for CoTPP, CuTPP, FeTPP, H₂TPP, NiTPP and ZnTPP.

Fig. 2 shows the CVs of H₂TPP, FeTPP, CoTPP, NiTPP, CuTPP and ZnTPP. All porphyrins have open circuit potentials around –0.3 V vs. Fc/Fc⁺ and in relation to this undergo 1–3 oxidations and 1–2 reductions in the solvent and potential window studied here. The redox transitions can occur either at the porphyrin ligand or at the metal center. Ligand-centered transitions will from here on be referred to as L^m/Lⁿ for shifts between total charges *m* and *n*, and metal-centered reactions will be referred to as M(p)/M(q) for transitions between metal oxidation state *p* and metal oxidation state *q*.

All compounds display the ligand-centered redox transitions L¹⁺/L⁰ and L²⁺/L¹⁺. H₂TPP, CoTPP, NiTPP and ZnTPP also undergo the L⁰/L^{1–} transition, whereas FeTPP only shows this transition in square wave voltammetry. For H₂TPP the π -radical anion to dianion transition L^{1–}/L^{2–} is visible in the CV, and for CuTPP and FeTPP it is visible in square wave voltammetry (see Fig. S1†). CoTPP and FeTPP also undergo the metal centered transitions M(II)/M(I) and M(III)/M(II). All observed redox reactions are electrochemically quasi-reversible except the L⁰/L^{1–} transition in CoTPP, which is irreversible. For this porphyrin, reoxidation from L^{1–} to L⁰ appears to occur in two steps, at a small shoulder at approximately –1.46 V vs. Fc/Fc⁺ and at the anodic peak at –0.79 V vs. Fc/Fc⁺. These features are not present when the negative potential cutoff is set positive of the L⁰/L^{1–} peak (see Fig. S2†).

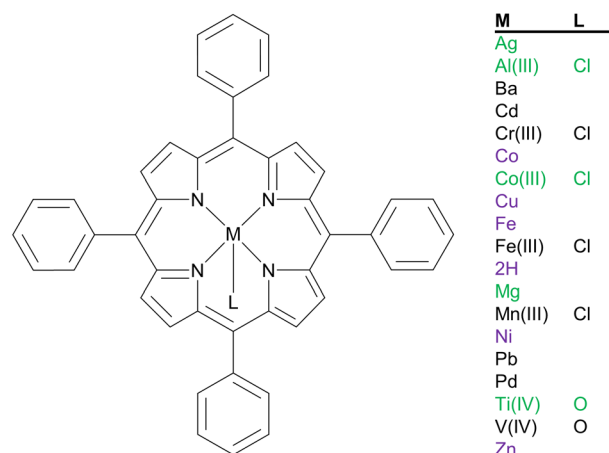


Fig. 1 Structures used in this work. For compounds marked in purple, both the electrochemical and catalytic properties were studied experimentally. For compounds marked in green, only the electrochemistry was evaluated. Compounds marked in black have only been studied computationally.



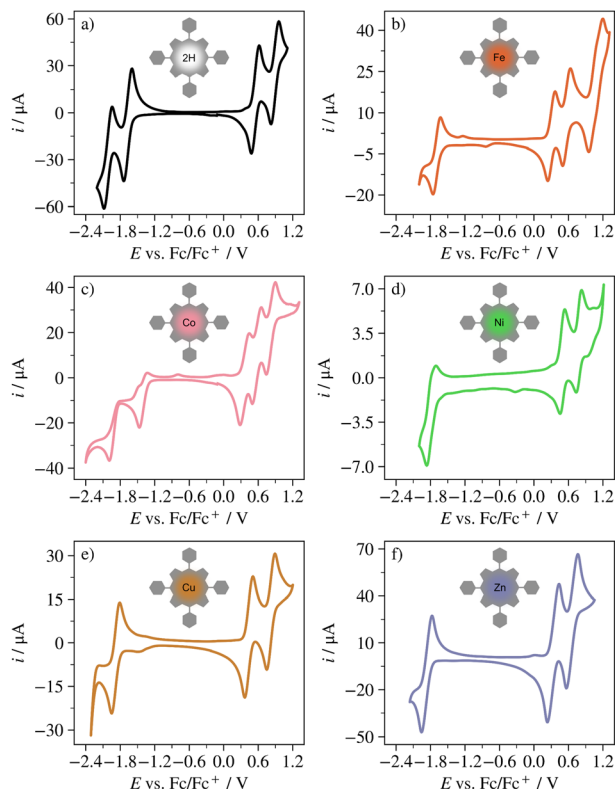


Fig. 2 CV of (a) H₂TPP, (b) FeTPP, (c) CoTPP, (d) NiTPP, (e) CuTPP, and (f) ZnTPP in 0.1 M TBAPF₆/DCM at 0.1 V s⁻¹.

NiTPP has previously been reported to be able to undergo the Ni(III)/Ni(II) transition,^{48–50} but this reaction occurs close to or even overlaps the L¹⁺/L⁰ transition, and in some reports it is not included.^{51–53} Under the reaction conditions used here (room temperature, 0.1 M TBAPF₆/DCM) the first redox conversion of NiTPP at oxidative potentials has been reported to correspond to the ligand-centered reaction L¹⁺/L⁰.⁵² This is also true for *e.g.* ((*p*-Cl)TPP)Ni⁵² and Ni(II) *meso*-(4-fluoro-2,6-dimethylphenyl)porphyrin (NiTFP)¹¹ studied under the same conditions. Given that the potential difference between the first redox transition at reductive potentials and the first transition at oxidative potentials is 2.31 V, in good agreement with both H₂TPP (2.2 V), ZnTPP (2.21 V) and CuTPP (2.32 V) for which no metal transitions occur, Ni is here assumed to remain in oxidation state +II. The redox chemistry of H₂TPP, FeTPP, CoTPP, NiTPP, CuTPP, and ZnTPP is summarized in Table 1. Further details, as well as the redox chemistry of AgTPP, Al(III)TPP, CoTPP in 0.1 TBAPF₆/dimethylformamide, Co(III)TPP and MgTPP can be found in Fig. S3, S4 and Table S1.†

The half-wave potentials of the porphyrin redox transitions (as for many other compounds) vary with conditions such as solvent, electrolyte and temperature. A thorough summary of previously reported half-wave potentials of a large number of tetraphenylporphyrins studied in various solvents can be found in Fig. S5, S6 and Table S2.† Except for the Fe(III)/Fe(II)

Table 1 Half-wave potentials of the observed redox transitions as measured by square wave voltammetry (V vs. Fc/Fc⁺, 0.1 M TBAPF₆/DCM, 15 Hz, 4 mV increment, 25 mV amplitude). *E*_{+/-} is the potential difference between the first half-wave potential at oxidative potentials, and the first half-wave potential at reductive potentials

Porphyrin	L ¹⁻ /L ²⁻	L ⁰ /L ¹⁻	M(III)/M(II)	M(III)/M(II)	L ¹⁺ /L ⁰	L ²⁺ /L ¹⁺	<i>E</i> _{+/-}
H ₂ TPP	-1.99	-1.64			0.56	0.90	2.20
FeTPP	-2.26	-2.07	-1.71	0.27	0.53	0.97	1.98
CoTPP		-1.91	-1.39	0.34	0.55	0.79	1.73
NiTPP		-1.82			0.49	0.78	2.31
CuTPP	-2.32	-1.88			0.44	0.82	2.32
ZnTPP		-1.85			0.36	0.66	2.21

conversion,^{48,49,54–57} all half-wave potentials measured in this work occur within the range of previously reported values.^{48–77}

Within the literature, minor disagreement remains regarding assignment of certain redox transitions. For example, the first redox transition at oxidative potentials in Al(III)TPP and AgTPP have been reported as both M(III)/M(II) and L¹⁺/L⁰, and some reports include the M(II)/M(I) transition of AgTPP and Cr(III)TPP whereas others do not. These discrepancies have not been further investigated here.

It should be noted that half-wave potentials in the literature are originally often reported against the standard calomel electrode (SCE), or on occasion against Ag/AgCl or the saturated sodium chloride calomel electrode (SSCE). The standard potentials for these reference electrodes vary with conditions, meaning that translation and comparison between them will likely introduce systematic errors. In this work, we have calibrated the SCE vs. the Fc/Fc⁺ scale by comparing literature half-wave potentials vs. SCE of H₂TPP (which are plentiful and in quite close agreement with each other across solvents and studies), with our own measured half-wave potentials of H₂TPP vs. Fc/Fc⁺. The average of the differences in half-wave potentials results in an approximate standard potential *E*_{SCE}⁰ = -0.48 V vs. Fc/Fc⁺, which falls well within the range of SCE standard potentials reported for various organic solvents.^{78–82} Given that *E*_{SSCE}⁰ = 0.244 V vs. SHE,⁸³ that *E*_{Ag/AgCl}⁰ = 0.209 vs. SHE⁸⁴ and that *E*_{SSCE}⁰ = 0.235 vs. SHE,⁸⁵ we have also arrived at *E*_{Ag/AgCl}⁰ = -0.515 vs. Fc/Fc⁺, and *E*_{SSCE}⁰ = -0.489 V vs. Fc/Fc⁺. These values should not be taken as universal, but only as a means to compare half-wave potentials on the same scale within this work.

From the CVs in Fig. 2, it is clear that the effect of the metal on the porphyrin redox chemistry in general manifests simply as a collective shift of all redox reactions towards more negative potentials compared to the half-wave potentials of the free base (the exception being FeTPP, for which the L²⁺/L¹⁺ occurs at slightly more positive potentials than for H₂TPP). This means that unless the metal itself is redox active, only its electron withdrawing abilities is affecting the redox chemistry of the ligand. Indeed, it has been showed that for a large number of octaethylporphyrins with metal centers remaining in oxidation state +II, the half-wave potential of the L¹⁺/L⁰ transition is proportional to the electronegativity of the metal



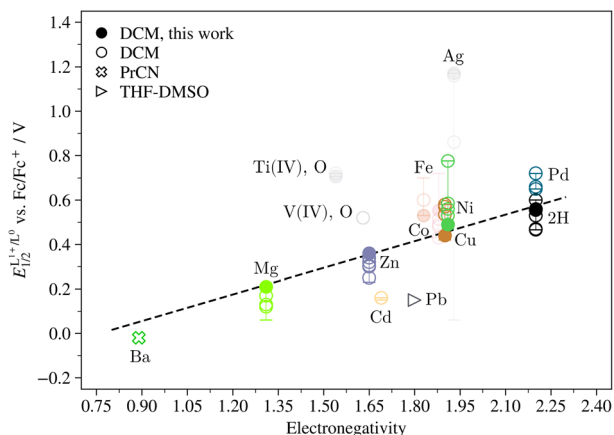


Fig. 3 Half-wave potential of the L^{1+}/L^0 transition as a function of metal (or hydrogen) electronegativity. Ti(IV), O and V(IV), O correspond to $O=Ti(IV)TPP$ and $O=V(IV)TPP$, respectively. Filled markers indicate values obtained experimentally in this study, open markers indicate values obtained from literature and the bars represent all literature values obtained in solvents other than DCM. Faded colors show which metals are not in the +II oxidation state.

center.⁸⁶ This holds true here as well, as shown in Fig. 3. It is also interesting to note that the difference in half-wave potentials between H_2TPP and $MTPP$ is larger for the L^0/L^{1-} transition than for the L^{1+}/L^0 transition. This indicates that the HOMO is more strongly affected by the presence of the metal than the LUMO.

4.2 Computational study

The catalytic cycle of the HER in porphyrins can be described as an overall four-step process involving two chemical protonations (P) and two reductive electron transfers (E). There are several possible reaction pathways, such as EEPP, EPEP, EPPE, PEEP, PEPE, and PEE (outlined in Scheme S1†). Which one is favored may depend on *e.g.* the proton source,^{14,87,88} the ligand structure and the metal center.⁸⁸ However, the EEPP^{13,87} and EPEP^{10,11,89} pathways dominate among previously reported mechanisms for porphyrins or porphyrin-like structures. As will be discussed in Section 5, these pathways are likely to be applicable in this study as well. During catalysis of the HER by tetraphenylporphyrins, a key intermediate should therefore be $H-MTPP$ with an overall charge of -1 (EEPP) or 0 (EPEP). Thus, the binding energy of H ($\Delta\epsilon_H$) can possibly be used as a descriptor of catalytic activity and as the foundation of a volcano plot for a series of $MTPP$ catalysts.

The binding energy of H relative a free $MTPP$ porphyrin is given by

$$\Delta\epsilon_H = \epsilon_{H-MTPP} - \epsilon_{MTPP} - \frac{1}{2}\epsilon_{H_2} \quad (1)$$

where ϵ_{H-MTPP} is the energy of the intermediate complex, ϵ_{MTPP} is the energy of the porphyrin and ϵ_{H_2} is the energy of a free hydrogen molecule. In this study, H will be assumed to bind to the metal center of metalated porphyrins (forming a

metal-hydrogen or metal-hydride complex), and to N on one of the pyrroles of the free base porphyrin. Other binding sites are not considered, as the purpose of this work is to evaluate the binding energy of hydrogen as a common denominator descriptor of catalytic behavior that does not require knowledge of the full reaction pathway. It should nonetheless be noted that there is a possibility that the ligand is involved in the protonation, for example through intramolecular charge transfer. Indeed, there is a change in Mulliken spin density on the ligand between $CuTPP$ and $H-CuTPP$ (see Fig. S8†). However, as Mulliken population analysis tends to be sensitive to the basis set and the changes in CM5 charge on the ligand and on the metal between $H-MTPP$ and $MTPP$ was small for all compounds used in the experimental study, the effects of intramolecular charge transfer were not further investigated here. Details on ligand charges and spin densities are available in Fig. S8–S14, Tables S4 and S5.†

The appropriateness of the level of theory of the computational study was confirmed by plotting the half-wave potential of the first redox transition at oxidative potentials against the calculated difference in free energy between $MTPP^0$ and $MTPP^+$, $\Delta G_{+/0}$. As can be seen in Fig. 4, the relationship is more or less linear (the linear fit was produced using the experimental half-wave potentials measured in this work). $AlTPP$, $CoTPP$, $CrTPP$, $MnTPP$ and literature half-wave potentials of $FeTPP$ do not fit in the trend (see Fig. S15†), possibly because the first oxidation occurs on the metal rather than on the ligand, and because this transition is associated with coordination of a Cl ion that is not represented here. $AgTPP$ also undergoes metal oxidation, but is not associated with any axial ligands and fits into the trend. All electronic energies, zero-point energies, thermal corrections and the multiplicities used here can be found in Table S6,† and the coordinates of

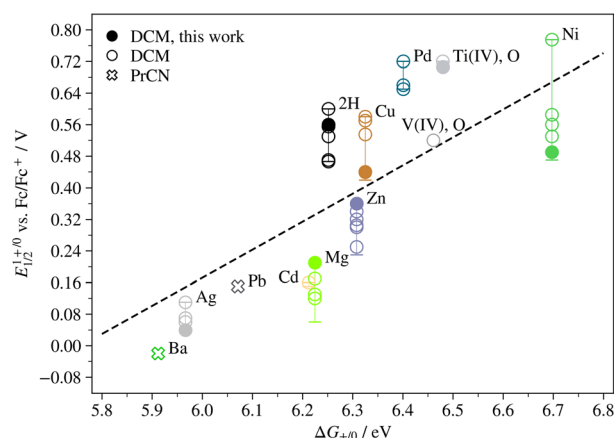
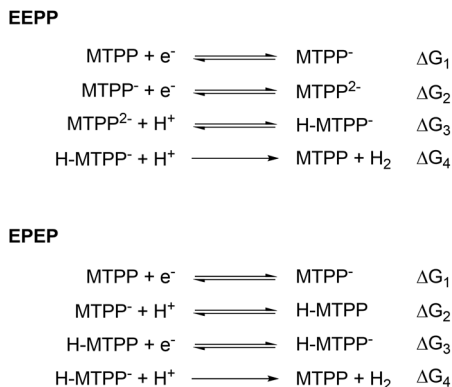


Fig. 4 Half-wave potential of the first redox transition at oxidative potentials ($M(III)/M(II)$ for $AgTPP$, L^{1+}/L^0 for all other compounds shown here) as a function of the difference in free energy between $MTPP^0$ and $MTPP^+$. Ti(IV), O corresponds to $O=Ti(IV)TPP$. Filled markers indicate values obtained experimentally in this study, open markers indicate values obtained from literature and the bars represent all literature values obtained in solvents other than DCM.





Scheme 1 The EEPP and EPEP reaction pathways for MTPP. Charges in this scheme are total for the entire complex, and reductions may occur either on the metal or on the ligand.

all optimized compounds are available in Appendix 1 of the ESI.†

As shown in Scheme 1, the four steps of the HER are associated with the changes in free energy ΔG_1 , ΔG_2 , ΔG_3 and ΔG_4 , respectively (note that metal- and ligand centered reductions are not distinguished):

As discussed above, the electron withdrawing ability of the metal center, *i.e.* the charge density ratio between the metal and the ligand, determines the ease of ligand reduction. It should also affect the ease of protonation, as protons will be more likely to adsorb onto a charge-depleted metal center. If linear free energy relations apply, the free energy change to produce the adsorbed state, ΔG_{H} , should be proportional to $\Delta G_1 = -nFE_{0/-}$, where $E_{0/-}$ is the formal potential (here assumed to be equal to the half-wave potential) of the first porphyrin reduction involved in the catalytic cycle. The free energy change of the adsorbed state is given by

$$\Delta G_{\text{H}} = \Delta \varepsilon_{\text{H}} + \Delta \varepsilon_{\text{ZPE}} - T\Delta S \quad (2)$$

where $\Delta \varepsilon_{\text{ZPE}}$ is the difference in zero-point energy between H-MTPP (or H-MTPP⁻), MTPP and $\frac{1}{2}\text{H}_2$, T is the temperature and ΔS is the change in entropy due to the adsorption.

Eqn (2) indicates that if ΔG_{H} is linearly related to ΔG_1 and thereby to $\Delta E_{0/-}$, so is $\Delta \varepsilon_{\text{H}}$. Indeed, for metals that are expected to remain in oxidation state +II, the binding energy is linearly related to the half-wave potential of the first ligand reduction (the linear fit was produced using the experimental half-wave potentials measured in this work). This is shown in Fig. 5a. As can be seen in Fig. 5b, there is also a linear trend between $\Delta \varepsilon_{\text{H}}$ and the half-wave potential of the first ligand oxidation. These results imply that the electron withdrawing ability of the central ion is the only determining parameter of the binding energy of H. They also imply that the binding energy of H agrees with the thermodynamics of the porphyrin redox reactions. This means that if the premise of the Sabatier principle holds true for porphyrins, *i.e.* that if their thermodynamic profiles are sufficient to predict their catalytic quality, the $\Delta \varepsilon_{\text{H}}$ should serve as a descriptor of catalytic activity for porphyrins just like it does for metal surface catalysts.

It should be pointed out that the binding energies in this work do not represent a single step in the free energy landscape (as they are assumed to do for catalysis of the HER on metal surfaces), but rather as a sum of two or more steps. In addition, binding energies in experimental settings depend on local reaction conditions such as electrode surface charge,¹⁵ and the free energy landscape at realistic operating conditions may not be flat.^{31–37,90} The binding energies given here should therefore not be considered quantitative. Nonetheless, a few detailed features should be mentioned: firstly, CoTPP, which undergoes the Co(II)/Co(I) transition before the L⁰/L¹⁻ transition, does not fit into the trend in Fig. 5a. Interestingly, the half-wave potential of the Fe(II)/Fe(I) transition does. As will be discussed in Section 5, the Fe(II)/Fe(I) transition partakes in catalysis of the HER, whereas the Co(II)/Co(I) transition does not. Secondly, the binding energies calculated here are all larger than 0, meaning that all compounds are expected to display weak binding towards H under ideal thermodynamic conditions. Lastly, the free base porphyrin fits into the trends for both reduction and oxidation. This strengthens the argument that the binding energy is in fact independent of the identity of the central metal ion. Rather, it is determined exclusively by the electron density on the ligand.

4.3 Catalysis

The catalytic ability towards the HER was evaluated experimentally for H₂TTP, FeTTP, CoTTP, NiTTP, CuTTP and ZnTTP. Fig. 6 shows the CVs of these porphyrins in 0.1 M TBAPF₆/DCM containing 0–10 equivalents of BzOH. All compounds show an increase in cathodic current paralleled by a decrease in reversibility of porphyrin redox chemistry with increasing acid concentration, both indications of catalytic activity towards the HER. The response of CuTTP strongly resembles that of Cu(II) *meso*-(4-fluoro-2,6-dimethylphenyl)porphyrin (CuTFP), which was proven efficient for catalysis of the HER under the same reaction conditions.¹¹ For FeTTP, adding BzOH to the solution results in a new quasireversible reaction at approximately −0.8 V vs. Fc/Fc⁺.‡

At 10 equivalents of BzOH, the catalytic waves of CoTTP and NiTTP approach the S-shape associated with scan rate independent, pure kinetic control of the catalytic reaction.^{11,91–94} For H₂TTP, FeTTP, NiTTP and ZnTTP, the catalytic wave is peak shaped, indicative of mass-transport limitations,^{11,91–94} or more rarely catalyst deactivation or blocking of the electrode surface.⁹⁵ Because the catalytic current never decreases with acid concentration for any of the porphyrins (see ref. 93), catalyst deactivation seems unlikely. Some electrode fouling was observed for CuTTP (see Fig. S16†), but as the adsorbed species did not give rise to any reduction waves and produced current magnitudes much smaller than those of the catalytic

‡ A possible explanation could have been that Fe leaves the ligand and interacts with BzOH. However, this scenario was ruled out by cycling Fe(III) trifluoromethanesulfonate under the same conditions as FeTTP. No new peaks occurred when BzOH was added to the solution, and the origin of the ones seen for FeTTP under catalytic conditions currently remains unexplained.



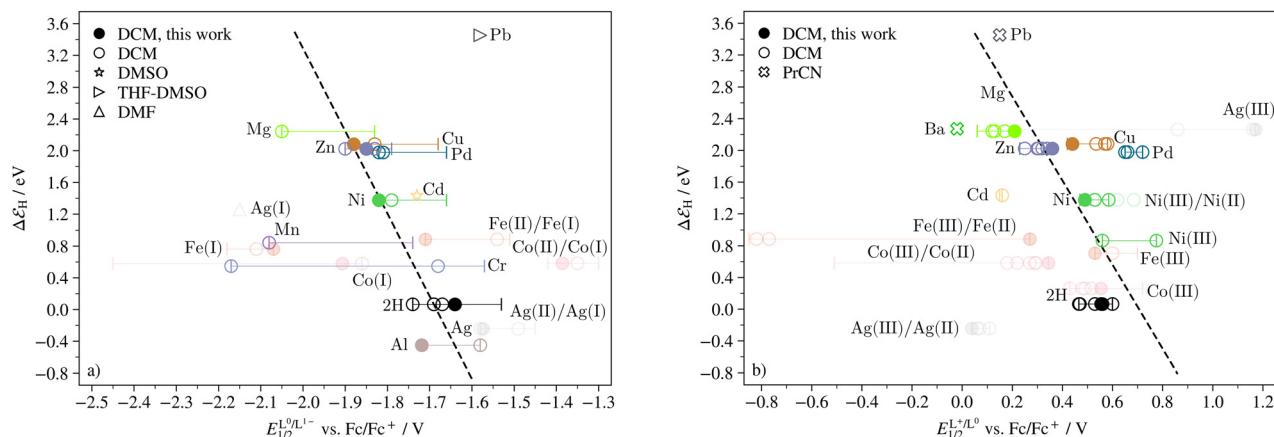


Fig. 5 Binding energy of H as a function of (a) the half-wave potential of the L^0/L^{1-} transition and (b) the L^{1+}/L^0 transition. Filled markers indicate values obtained experimentally in this study, open markers indicate values obtained from literature and the bars represent all literature values obtained in solvents other than DCM. Faded colors show which metals are not in the +II oxidation state. The $M(II)/M(I)$ transition for AgTPP, CoTPP and FeTPP, and the $M(III)/M(II)$ transition for AgTPP, CoTPP, FeTPP, and NiTPP (the latter not seen in all studies), are included for reference. The binding energy for Ag(II)TPP is plotted against both the half-wave potential of what is here assumed to be the $Ag(II)/Ag(I)$ transition, and against what has been previously been reported as the L^0/L^{1-} transition.⁶² Similarly, the binding energy for Ni(II)TPP is plotted against both the half-wave potential of what is here assumed to be the L^{1+}/L^0 transition, and against what has elsewhere been reported as the $Ni(III)/Ni(II)$ transition.^{48–50}

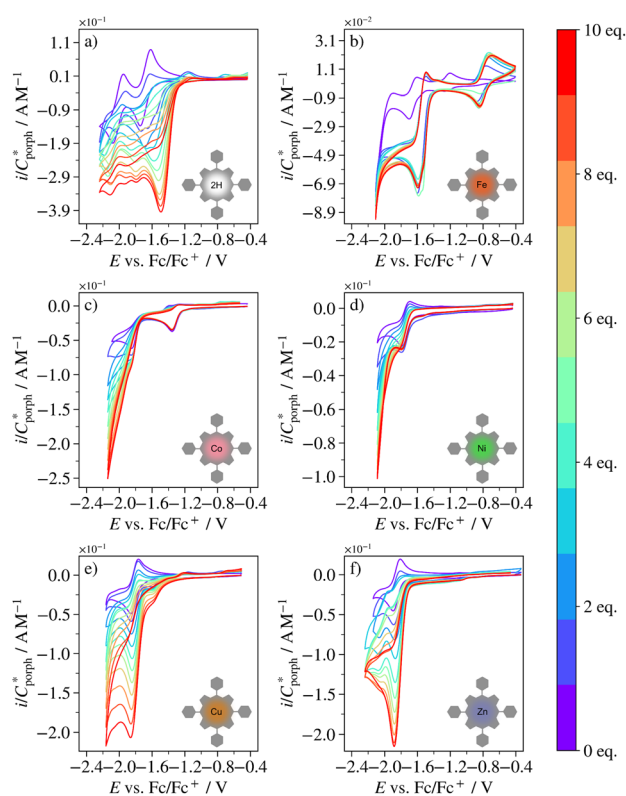


Fig. 6 CVs at 0.1 V s^{-1} of (a) H_2TPP , (b) $FeTPP$, (c) $CoTPP$, (d) $NiTPP$, (e) $CuTPP$ and (f) $ZnTPP$ in $0.1 \text{ M TBAPF}_6/\text{DCM}$ containing 0–10 equivalents of $BzOH$ (indicated by colorbar).

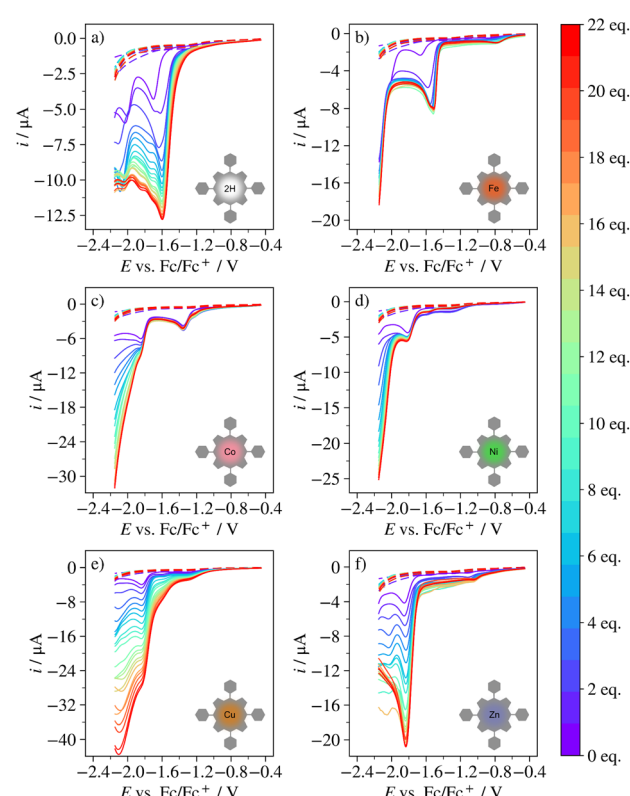


Fig. 7 LSV at 0.1 V s^{-1} of (a) H_2TPP , (b) $FeTPP$, (c) $CoTPP$, (d) $NiTPP$, (e) $CuTPP$ and (f) $ZnTPP$ in $0.1 \text{ M TBAPF}_6/\text{DCM}$ containing 0–22 equivalents of $BzOH$ (indicated by colorbar).

reaction, inhibition of the catalytic mechanism through blocking of the electrode surface was deemed negligible for all porphyrins.

4.3.1 Catalytic mechanism. The appearance of the catalytic wave varies widely between the porphyrins. This is clearly visible in Fig. 7, showing LSVs of all porphyrins in the presence of 0–22 equivalents of Benzoic acid. For H_2TPP , the cata-



lytic reaction begins in connection to the L^0/L^{1-} redox transition. Between 1 and 4 equivalents of acid, the cathodic peak potential of this transition shifts towards more positive values, and thereafter remains constant. This suggests that the first two steps of the catalytic mechanism involve one reduction and one protonation, where the proton is strongly bound to the porphyrin, *i.e.* that the reduced and protonated intermediate is stable. Such a process would cause the L^0 state of the porphyrin to be stabilized relative the L^{1-} state, leading to the observed potential shift. This observation also agrees with trend in the computational data, which indicates that the free base has a stronger binding energy of H than the other porphyrins studied here.

FeTPP behaves similarly to H_2 TPP between 1 and 5 equivalents of acid, in that the catalytic wave occurs in connection to the first reduction wave of the porphyrin (here the $Fe(II)/Fe(I)$ transition) and in that the cathodic peak potential of this wave shifts towards less negative potentials. Like for H_2 TPP, the first two steps of the catalytic mechanism for FeTPP are thus likely to involve one reduction and one protonation, where the intermediate is stable. However, above 5 equivalents, a current plateau occurs where the reaction is independent of both acid concentration and potential until the potential is sufficiently negative that a second reductive wave occurs (likely part of the L^0/L^{1-} transition, given that this occurs approximately 0.4 V below the $Fe(II)/Fe(I)$ transition). The behavior of FeTPP indicates that at more than 5 equivalents of acid, all catalytic units at the electrode surfaces are occupied by a bound proton and the catalytic mechanism cannot continue until a second reduction occurs and/or the driving force is increased dramatically. In fact, it has been shown before that the hydrogen evolution mechanism may change from EPEP to EEP during a cathodic potential sweep for certain Ni-based catalytic complexes.^{96,97} It should be noted that the catalytic current response observed for FeTPP here is different from that observed for $Fe(III)TPP$ in 0.1 M tetraethylammonium perchlorate/dimethylformamide in the presence of triethylamine hydrochlorate.¹²

For CoTPP, no catalysis occurs at the first reduction wave of the porphyrin, corresponding to the $Co(II)/Co(I)$ transition. Rather, the catalytic wave begins only at the L^0/L^{1-} transition, where it approaches the characteristic S-shape at 4 equivalents of acid. No potential shift is observed, but the catalytic wave is convoluted by what may be the L^2-/L^{1-} transition. The half-wave potential of this transition has not been reported, but the background current generated by glassy carbon is negligible in the relevant potential region. Given that the metal peak is not involved in catalysis and that the ligand peak is unaffected in terms of potential, the first step of the catalytic mechanism may be reduction of the ligand, or protonation yielding a short-lived intermediate. CoTPP has previously been reported to catalyze the HER through the EEP mechanism by undergoing the $Co(II)/Co(I)$ transition followed by the $Co(I)/Co(0)$ transition.¹³

The LSVs of NiTPP are similar to those of both FeTPP and CoTPP. There is a slight increase in the cathodic current at the

L^0/L^{1-} transition, but at 3 equivalents of acid, a short current plateau appears. Like for FeTPP, this may indicate that the first two steps of catalysis involve one reduction and one protonation, producing an intermediate complex that is stable enough to impede catalysis until the driving force has been further increased. However, no potential shift is seen for NiTPP, suggesting the intermediate is slightly less stabilized by protonation for NiTPP than for FeTPP. This is corroborated by the fact that the second part of the catalytic wave occurs at less negative potentials for NiTPP than for FeTPP, and by the computational result indicating that the binding energy of H is stronger for FeTPP than for NiTPP.

For CuTPP and ZnTPP, the catalytic wave occurs at the L^0/L^{1-} transition, without peak shift and current plateau. This indicates that the first steps of the mechanism involve one reduction and possibly a protonation leading to a short-lived intermediate. At 11 equivalents, the current response of CuTPP approaches the canonical S-shape. For ZnTPP, the current response remains peak-shaped throughout all acid concentrations, but stops increasing in magnitude at 17 equivalents. Thus, the ZnTPP mechanism eventually “saturates” similarly to that of FeTPP and NiTPP. These observations suggest that the intermediate is slightly more stable for ZnTPP than for CuTPP, which agrees with their relative $\Delta\epsilon_H$.

From the CVs and LSVs, it is clear that the catalytic electrochemistry is different for all porphyrins. H_2 TPP and FeTPP produce substantial potential shifts, FeTPP and NiTPP display current plateaus after a certain amount of acid has been added, and the metal reduction of FeTPP is involved in catalysis but the one of CoTPP is not. CuTPP and ZnTPP show no potential shifts or sluggish initial steps of the catalytic mechanism, but as opposed to the one of CuTPP, the mechanism of ZnTPP is limited by mass-transport. The behaviors of the porphyrins can be linked to their binding energies, but only to the extent that the binding energies agree somewhat with the apparent sluggishness of the initial steps of the catalytic reaction and/or the stability of intermediates formed during these steps. The binding energy cannot fully explain the differences in catalytic mechanisms between the porphyrins. For example, both H_2 TPP and FeTPP display large potential shifts and have stronger binding than *e.g.* CuTPP and ZnTPP, but only FeTPP reaches “saturation”. Furthermore, CoTPP, FeTPP, and NiTPP all have stronger binding than CuTPP and ZnTPP and all require two reductions to be catalytic at most acid concentrations. However, $\Delta\epsilon_H$ does not relate to the length of the current plateau, the number of acid equivalents needed to reach the plateau, or the potential at which the second reduction wave occurs.

The porphyrins behave similarly in that the first two steps of the catalytic mechanism seem to involve at least one reduction. Fig. 8 shows the UV/vis absorption spectra of all porphyrins in the presence of 0 and 22 equivalents of BzOH. The Soret and Q band peaks of the two spectra overlap exactly for all porphyrins, and the spectra with acid present are simply an overlay of the respective porphyrin spectrum and the acid



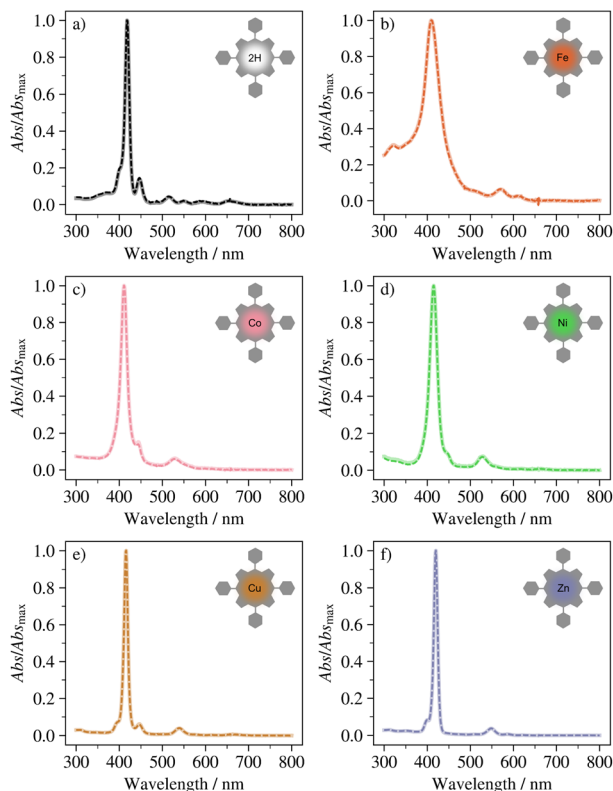


Fig. 8 UV/vis absorption spectra of each porphyrin in the presence of 0 (full line) and 22 (dashed line) equivalents of BzOH. All spectra were recorded in DCM and normalized by the maximum absorption Abs_{max} .

spectrum (the spectrum of the same amount of acid without any porphyrin present is shown in Fig. S17†). These results show that no protonation occurs when no potential is applied, which together with the above observations suggests that reduction is the first step of the catalytic mechanism for all porphyrins. This is in agreement with previous studies on similar systems.^{10,11,13,87,89} However, it should be noted that protonation may still occur at potentials less negative than the half-wave potential of any porphyrin reduction, but more negative than the open circuit potential at which the porphyrin is in its L^0 state.

4.3.2 Reaction kinetics: potential dependence of the catalytic current. To evaluate the efficiency of a catalyst, the reaction kinetics must be quantified. This is notoriously difficult to do with accuracy for multistep, multielectron processes, especially when the catalytic mechanism is not fully known. The reaction mechanism for porphyrins catalyzing the HER is often reported as EEPP or EPEP, but the reaction may involve more steps, the steps may occur in another order, and the order may vary between the different porphyrins. Consequently, any expression of reaction rates or rate constants will be elaborate (an example is given in Section 3.3.1. of the ESI†), may differ between porphyrins, will not allow distinguishing between the rates of the different steps, and anything less than ideal experimental conditions will not allow for

steady state approximations. All rates and rate constants will be global, offering no information of local variations. However, it may still be possible to obtain qualitative trends.

A first qualitative evaluation of the reaction rates and limitations can be achieved by studying how the current varies with the concentration of acid at different potentials. The two protonations will be dependent on the concentration of protons (here equaled to the concentration of acid, as BzOH is not expected to dissociate in DCM) but not on the potential, and the opposite will be true for the two reductions. If the current is *dependent* on the acid concentration at a given potential, protonation and reduction proceed at similar rates; if the current is *independent* of the acid concentration, the electrochemistry and release of the product must be limiting as increasing the concentration of protons does not increase the turnover frequency. Thus, plotting the current as a function of acid concentrations at different overpotentials should show which redox event is triggering for the catalytic cycle and to what extent this redox event is limiting on the reaction rate.

In heterogeneous catalysis, the current density at a given overpotential is a common measure of catalytic activity.⁹¹ However, in molecular catalysis, the concept of overpotential is not straightforward. The overpotential describes the additional potential required to drive a reaction at a certain rate compared to the standard potential of the reaction,^{91,98} but a molecular catalytic cycle is usually nonexistent until triggered by *e.g.* a redox event, after which the rate of the reaction is determined by a series of possibly both potential-independent chemical steps and potential-dependent electrochemical events.⁹¹ In addition, to determine the true overpotential, the standard potential of the reaction to be catalyzed (here the HER) must be known under the relevant reaction conditions. The standard potential and pK_a of the HER in DCM using BzOH as a proton source have not yet been well defined. These values could be determined through *e.g.* open circuit potential measurements,⁹⁹ but an accurate description of the behavior of BzOH in DCM is outside the scope of this study. For these reasons, we instead of true overpotentials and the standard potential of the HER, use a range of reference potentials, E_{ref} , to represent different driving forces of the HER.

Fig. 9 shows how the current generated by the catalytic reaction (estimated by subtracting the current response of the porphyrin without any acid present, i_0 , from all other currents) as a function of acid concentration at several E_{ref} . At small E_{ref} magnitudes, the currents are small and vary noisily over the acid concentrations. At $E_{ref} = -1.6$ V vs. Fc/Fc^+ , H_2TPP and $FeTPP$ are most active. This is expected, considering that this reference potential is close to the half-wave potentials of the first reduction – the triggering event for catalysis – of these porphyrins. No activity is seen for $CoTPP$ at this potential, which in agreement with the LSVs suggests that the reduction from $Co(II)$ to $Co(I)$ is not enough, or does not occur at negative enough potentials, to initiate the catalytic cycle. The activity of $CuTPP$, $NiTPP$ and $ZnTPP$ at $E_{ref} = -1.6$ V vs. Fc/Fc^+ is also negligible. This is also expected, as the first reduction wave of these porphyrins all occur below -1.8 V vs. Fc/Fc^+ , and as dis-



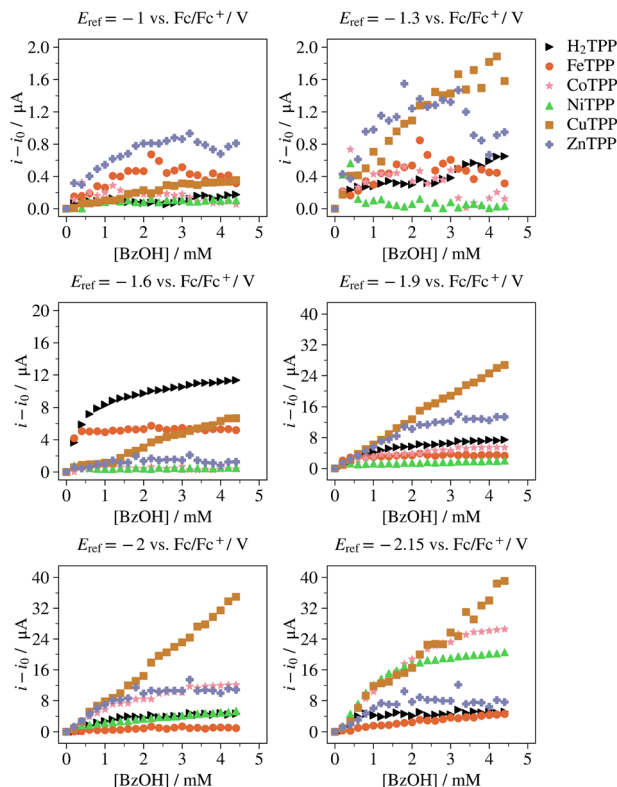


Fig. 9 Current responses as an estimate of reaction rates at different potentials, extracted from the results in Fig. 7.

cussed above, a single reduction of NiTPP is not enough to drive the catalytic reaction at high acid concentrations. At $E_{\text{ref}} = -1.9$ V vs. Fc/Fc^+ , the activity of CuTPP, CoTPP, NiTPP and ZnTPP increases, with the current being consistency highest for CuTPP. For H_2TPP and FeTPP, the catalytic current reaches a plateau, or semi-plateau, at all reference potentials. This means that the catalytic cycle becomes independent of the acid concentration, suggesting that the catalytic turnover is too slow to accommodate more than a few equivalents of protons. From -2 V vs. Fc/Fc^+ , the current for all porphyrins except CuTPP reaches a plateau. This indicates that the catalytic turnover for CuTPP is faster than for the other porphyrins, such that regardless of acid concentration, there is no “saturation” or delay in conversion.

From these rates and their variation with the acid concentration, reaction orders in $[\text{BzOH}]$ can be obtained. As shown in Fig. S18–S23,† the rate of the catalytic reaction for all porphyrins is linearly dependent on $[\text{BzOH}]$ when $[\text{BzOH}]$ is small, suggesting that the catalytic reaction is first order in acid concentration. At larger acid concentrations, all porphyrins approach zeroth order in acid concentration except CuTPP. It should be noted that the breaking point between first and zeroth order is not well defined for CoTPP, and to some extent NiTPP (see Fig. S20 and S21†). Rather, there is a breaking “arc” between the two regions. This suggests that the reaction cannot be accurately described by a steady-state approximation for these compounds.

As shown in Fig. S25,† the catalytic reaction is also first order in porphyrin concentration, indicating that the catalytic reaction is monomolecular. This means that approximate, global rate constants, k_{obs} , can be obtained from:

$$\frac{d[\text{H}_2]}{dt} = k_{\text{obs}}[\text{MTPP}][\text{H}^+] \quad (3)$$

where $k_{\text{obs}}[\text{MTPP}]$ corresponds to the slope of a linear fit of the reaction rate vs. $[\text{BzOH}]$. Here, it is important to note that if the mechanisms vary between the porphyrins, the stepwise rate constants contained within k_{obs} will also vary, and the k_{obs} will not be comparable between the different porphyrins. Indeed, when the rates as well as k_{obs} obtained in the above manner are plotted against the binding energy of H, there are no clear trends (see Fig. S26 and Fig. S27†) and k_{obs} cannot be used to create an experimentally relevant volcano plot. However, at the most negative E_{ref} , the reaction rates scale with the half-wave potentials of the first reduction wave involved in the catalytic cycle (see Fig. S28†). This is reasonable if the first step of catalytic cycle is reduction for all porphyrins, because if catalysis is triggered at a more negative potential, the catalytic unit will contain more energy to drive the catalytic reaction and the rate of the reaction will be higher.

The potential dependence of the catalytic current and the rate of the catalytic reaction can also be estimated by applying the Butler–Volmer equation, which governs the current at the electrode surface for an n electron process:¹⁰⁰

$$i = F A k^{\circ} \left(C_{\text{O}}(0, t) \exp \left[-\frac{\alpha n F}{RT} (E - E^{\circ}) \right] - C_{\text{R}}(0, t) \times \exp \left[\frac{(1 - \alpha) n F}{RT} (E - E^{\circ}) \right] \right) \quad (4)$$

Here, F is the Faraday constant, A is the electrode area, k° is the standard rate constant, α is the charge transfer coefficient, R is the gas constant, T is the temperature, E is the potential, E° is the formal potential of the reaction (here E_{ref} is used instead), and $C_{\text{O}}(0, t)$ and $C_{\text{R}}(0, t)$ are the concentrations at the electrode surface of the oxidized and reduced species, respectively.

Assuming that the back reaction is negligible, which should be true at high driving forces, the equation simplifies to the first term – the Tafel equation – and a rate can be obtained from a linear region of the logarithm of the current. Again, the current response of the porphyrin without any acid present can be subtracted to clarify the catalytic current response:

$$\ln(i - i_0) = \ln(F A k^{\circ}) - \frac{\alpha n F}{RT} (E - E_{\text{ref}}) \quad (5)$$

where $k^{\circ} = k^{\circ} C_{\text{O}}(0, t) = k^{\circ} C_{\text{cat}}(0, t)$ represents the reaction rate with respect to E_{ref} . The concentration of the catalyst at the electrode surface, $C_{\text{cat}}(0, t)$, is absorbed into the rate as it is not known. However, at high driving forces, it should be approximately equal to the bulk concentration. This method for evaluating the reaction kinetics is similar to the first one, barring



the fact that the rate here comes from an average of the current over a potential range, rather than being equal to the current at a specific potential. The Tafel fits used here are shown in Fig. S29–S34.† The rate behaviors are similar to those discussed above, but less well-defined as only a small portion of the logarithmic plots are close to linear and the approximation of the rate therefore will be further away from the truth the further away E_{ref} is from the relevant potential region. It is no surprise that the complexity of the mechanisms discussed here are not well described by idealized Tafel equations. As for the rates discussed above, there are no clear trends in reaction rates obtained in this manner with binding energies of H (see Fig. S36†), and an experimentally applicable volcano plot cannot be produced.

4.3.3 Reaction kinetics: current plateau analysis and catalytic Tafel plots. For an ideal, S-shaped catalytic wave, the plateau current can be used to obtain quantitative information about the reaction rate. For a catalytic reaction involving n heterogeneous electron transfers per catalyst and n' equivalents of catalyst per catalytic cycle, the plateau current is given by⁹¹

$$i_{\text{pl}} = nFAC_{\text{cat}}^* \sqrt{Dn'k_{\text{obs}}} \quad (6)$$

where C_{cat}^* is the bulk concentration of the catalyst, D is the diffusion coefficient and k_{obs} is the observed global rate constant of the rate limiting step or a convolution of elementary steps^{94,101} (note that this rate constant will still vary with acid concentration). For an EEP or EPEP process where both electron transfers occur at the electrode surface, $k_{\text{obs}} = k_1 C_s^*$, where C_s^* is the bulk concentration of substrate and k_1 is the rate constant associated with the first chemical step after the initial reduction(s). If this step is rate-limiting, k_1 can be taken as the rate constant of the catalytic reaction, k_{cat} .

The plateau current can be normalized by the peak current of a non-catalytic reaction, which is given by the Randles-Sevcik equation:^{91,92,101}

$$i_{0,\text{p}} = 0.4463n''FAC_{\text{cat}}^* \sqrt{\frac{n''FvD}{RT}} \quad (7)$$

where n'' is the number of electrons involved in the non-catalytic reaction and v is the scan rate. Here, $n = 2$ and $n' = n'' = 1$, yielding:

$$\frac{i_{\text{pl}}}{i_{0,\text{p}}} = \frac{n}{0.4463n''} \sqrt{\frac{n'RTk_{\text{obs}}}{n''Fv}} = 4.48 \sqrt{\frac{RTk_{\text{obs}}}{Fv}} \quad (8)$$

Eqn (8) and the analysis that lead to it apply when the catalytic wave is S-shaped, *i.e.* when the plateau current is independent of the scan rate and substrate diffusion. As this is not the case for all porphyrins studied here, this method can at best produce qualitative results, in which i_{pl} is approximated as the catalytic peak current.

Fig. 10(top) shows k_{obs} as a function of $[\text{BzOH}]$ for all porphyrins. These results reflect the observation that for FeTPP and NiTPP a single reduction is only sufficient to convert a small amount of substrate. CuTPP again appears to be able to

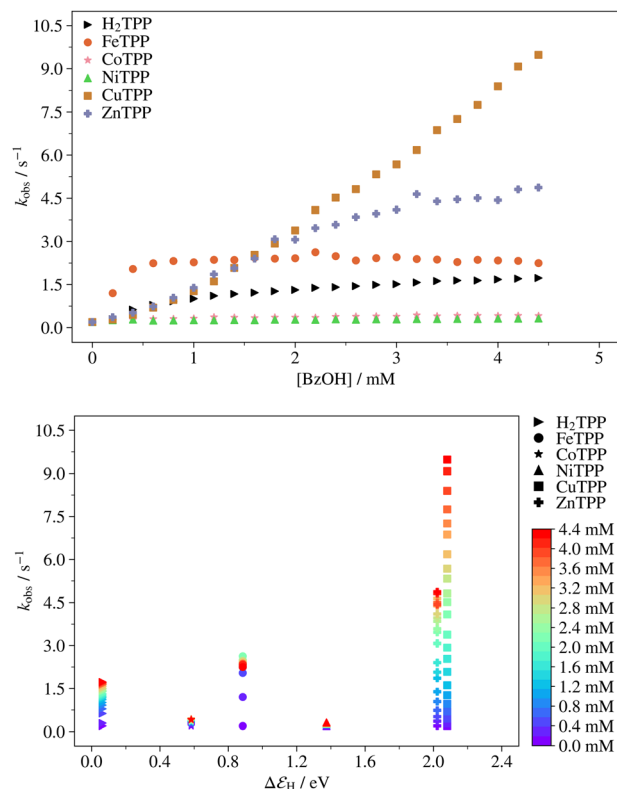


Fig. 10 Top: k_{obs} obtained from plateau current analysis of the results in Fig. 7. Bottom: k_{obs} obtained from current plateau analysis of the results in Fig. 7 vs. the binding energy of H. The colorbar indicates $[\text{BzOH}]$.

convert the largest amount, followed by ZnTPP. For H_2TPP , only a small increase in rate is observed for $[\text{BzOH}] > 1$ mM. The rate constants obtained through current plateau analysis agree with the calculated binding energies of H for H_2TPP , FeTPP, ZnTPP and CuTPP; H_2TPP has the strongest binding and is the slowest of these four porphyrins, and CuTPP shows the weakest binding and the fastest conversion. However, NiTPP and CoTPP do not fit into the trend and the correlation cannot be considered deterministic (see Fig. 10, bottom).

Like in the previous sections, k_{obs} obtained through current plateau analysis was used to calculate the reaction order in terms of $[\text{BzOH}]$. These results are shown in Fig. S38–S40.† Here, the catalytic reaction appears to be second order in $[\text{BzOH}]$ for CuTPP, at low concentrations for ZnTPP, and possibly at low concentrations of H_2TPP . However, in this analysis, the catalytic response was divided rather than subtracted by i_0 . At low concentrations of acid, the pure porphyrin response still constitutes a notable portion of the current response (visible especially for ZnTPP and CuTPP, where the cathodic current increases but does not change shape compared to i_0 at low acid concentrations). This means that both the catalyst concentration at the electrode surface and the acid concentration influence k_{obs} . Given that the catalyst concentration at the electrode surface is likely not equal to the bulk concentration of porphyrin, as there is evidence of substrate con-



sumption at low concentrations of acid, the result may be an apparent second order reaction with respect to [BzOH].

The rate constants obtained from current plateau analysis can be used to produce a so-called catalytic Tafel plot, where the turnover frequency (TOF) of the catalytic reaction is plotted as a function of the overpotential.⁹⁴ For a Nernstian process, the TOF is given by^{91,94}

$$\text{TOF} = \frac{\text{TOF}_{\max}}{1 + \exp\left[\frac{F}{RT}(E'^{\circ} - E_{\text{cat}/2})\right]} \exp\left[-\frac{F}{RT}\eta\right] \quad (9)$$

where the maximum turnover frequency $\text{TOF}_{\max} = k_{\text{obs}}$, E'° is the formal potential of the reaction to be catalyzed (here replaced by E_{ref}), η is the overpotential relative to E'° and $E_{\text{cat}/2}$ is the half-wave potential of the steady-state catalytic wave. It is often defined as the potential at which the catalytic current reaches half of its maximum value, but under pure kinetic conditions, $E_{\text{cat}/2}$ is assumed to be equal to the formal potential of the catalytic couple. Here, $E_{\text{cat}/2} = E_{0/-}$ will be used to obtain a qualitative comparison of turnover frequencies. Fig. 11 shows the catalytic Tafel plots for all porphyrins, for a range of different E_{ref} , using $\eta = E_{\text{ref}} - E$. At low overpotentials, H_2TPP and FeTPP have the highest TOF. This is expected, given that the onset of catalysis occurs at less negative potentials for H_2TPP and FeTPP than for the other porphyrins. At larger overpotentials for $E_{\text{ref}} < -1$ V vs. Fc/Fc^+ , the TOF of CuTPP and ZnTPP surpass those of H_2TPP and FeTPP , emphasizing the

importance of reaction conditions and kinetics when evaluating a catalyst.

4.3.4 Reaction kinetics: foot-of-the-wave analysis. For non-ideal catalytic responses, where the scan rate dependence of k_{obs} cannot be surpassed, foot-of-the-wave analysis (FOWA) can be used to evaluate reaction kinetics.^{94,95} The method is based on the idea that if a catalytic wave is misshaped by *e.g.* substrate consumption, the foot of the wave may still adhere to more purely kinetic conditions. Instead of approximating the plateau current as the peak current, an idealized plateau current is obtained from equations specific to the reaction mechanism at hand (this, of course, requires that the reaction mechanism is known). For an EEP or EPEP mechanism where both electron transfers are heterogeneous and the first chemical step is rate limiting, the current at the foot of the wave is given by^{91,94}

$$i = \frac{i_{\text{pl}}}{1 + \exp\left[\frac{F}{RT}(E - E_{\text{cat}/2})\right]} \quad (10)$$

A so called FOWA plot is obtained by plotting $i/i_{0,p}$ vs. the denominator of eqn (10). The rate constant can then be calculated from the slope of the linear section of this plot, which should occur close to the origin. This section corresponds to an idealization of the current response, and the resulting rate constant represents the rate of the reaction had the current response been ideal.

For a non-ideal catalytic response, the determination of $E_{\text{cat}/2}$ is not straightforward. Because the plateau current is not reached, it is not possible to exactly identify the potential of $i_{\text{pl}}/2$. However, in the ideal case, this potential should occur at or close to the inflection point of the catalytic wave.⁹⁸ As noted above, it can also be approximated as the formal potential of the catalyst couple. For the porphyrins discussed here, both $E_{\text{cat}/2}$ and the potential range of the foot of the wave are difficult to define. For example, for H_2TPP and FeTPP , the potential of the catalytic peak changes with concentration of the acid, and CoTPP has a reduction peak that is catalytically inactive before the actual catalytic peak. For this reason, we have performed the analysis using several different approximations of $E_{\text{cat}/2}$ and extensions of the linear range in the FOWA plot. Details are available in the ESI, Section 3.7.1.† No relevant comparison between the porphyrins could be made using any of the approximations, and no trends could be seen between k_{obs} obtained from FOWA and $\Delta\epsilon_{\text{H}}$. This is in line with all above discussions, which show that quantitative kinetic information and an experimentally relevant volcano plot are difficult to obtain, especially when the exact reaction mechanism is unknown.

In fact, none of the kinetic information obtained here is useful for anything else than a qualitative comparison between the porphyrins. The electrochemistry and thermodynamics of the compounds can be explained and follow simple trends, such as the dependence of the ligand redox chemistry and $\Delta\epsilon_{\text{H}}$ on the electronegativity of the metal center, and the thermodynamic linear scaling relation between steps in the catalytic

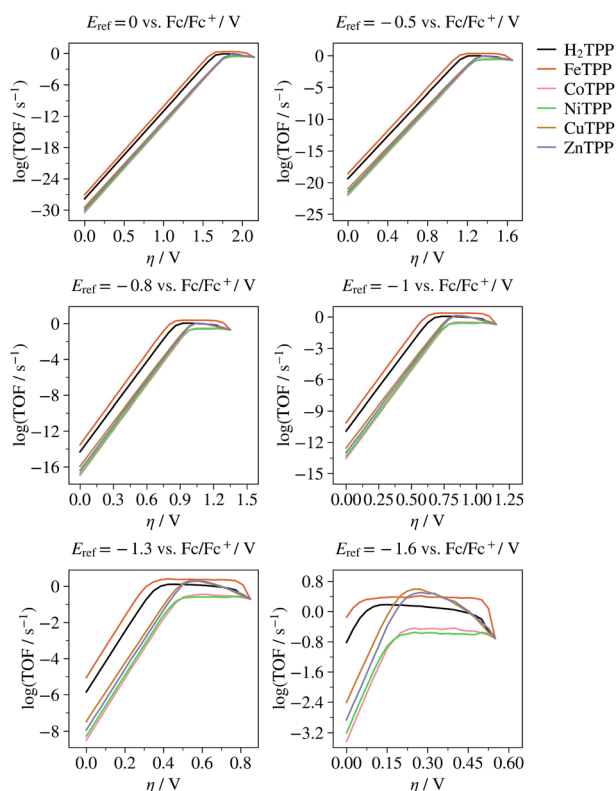


Fig. 11 Catalytic Tafel plots obtained through current plateau analysis of the results in Fig. 7.



cycle. However, as soon as kinetics are factored into the catalytic quality, the simple relations break down and there is no clear predictive power to the thermodynamic properties of the systems. An appropriate descriptor of catalytic efficiency likely must include effects of diffusion, the electronic double layer and/or interactions between the catalysts and the electrode surface. This work shows that tuning of catalytic properties through predictive models such as volcano plots and thermodynamic linear scaling relations cannot be applied without deep experimental knowledge of the systems.

5 Conclusion

We have reinvestigated the redox chemistry of a large number of tetraphenylporphyrins and found that reported values of half-wave potentials vary between different studies and experimental conditions. Nonetheless, we have seen that the redox chemistry of the porphyrins studied experimentally in this work fall within the range of previously reported behaviors, and that the half-wave potentials of ligand redox conversions are determined only by the electronegativity of the central metal ion.

We have also constructed a computational model to evaluate the applicability of Sabatier's principle and the volcano plot concept to experimental molecular catalysis of the HER. We have found that the binding energy of H correlates to the redox chemistry of the porphyrins, in that higher electronegativity of the central metal ion leads to stronger binding. This means that only the electron density of the ligand determines the longevity of what should be a key intermediate of the catalytic reaction, and that the first steps of the reaction are correlated by a linear scaling relation. However, under catalytic conditions the porphyrins studied here all behave differently and the catalytic mechanism cannot be clearly related to the redox chemistry of the porphyrins. Consequently, thermodynamics are not enough to accurately predict which molecular catalyst in a series will be most efficient and the binding energy of hydrogen is not a sufficient descriptor of experimental molecular catalysis.

The varying behaviors of the porphyrins under catalytic conditions also impedes proper comparison of kinetic parameters describing their catalytic cycles. All methods commonly applied to evaluate kinetic parameters of molecular catalysts, such as current plateau analysis or FOWA, require a high degree of ideality in the current responses and knowledge of the catalytic mechanism to be quantitatively accurate. For the series of porphyrins studied here, none of these methods could be used for more than a qualitative comparison of catalytic behavior. In summary, molecular design – often used as a selling point for molecular catalysis – is not straightforward. Construction of molecular volcano plots that are relevant to experimental operating conditions, prediction of catalytic properties and mechanisms, and fair comparison within a molecular series therefore remain strenuous tasks. Until reliable models have been developed, computational catalysis should

focus on reaction mechanisms rather than thermodynamic data and scaling relations.

Author contributions

Felicia Zaar: methodology, formal analysis, investigation, writing – original draft, visualization. Moyses Araujo: conceptualization, methodology, writing – review & editing. Rikard Emanuelsson: conceptualization, writing – review & editing, supervision. Maria Strømme: resources, writing – review & editing, funding acquisition. Martin Sjödin: conceptualization, methodology, resources, writing – review & editing, supervision, project administration, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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