ELECTROCHEMISTRY

Combining scaling relationships overcomes rate versus overpotential trade-offs in O₂ molecular electrocatalysis

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The development of advanced chemical-to-electrical energy conversions requires fast and efficient electrocatalysis of multielectron/multiproton reactions, such as the oxygen reduction reaction (ORR). Using molecular catalysts, correlations between the reaction rate and energy efficiency have recently been identified. Improved catalysis requires circumventing the rate versus overpotential trade-offs implied by such “scaling relationships.” Described here is an ORR system—using a soluble iron porphyrin and weak acids—with the best reported combination of rate and efficiency for a soluble ORR catalyst. This advance is achieved not by “breaking” scaling relationships but rather by combining two of them. Key to this behavior is a polycationic ligand, which enhances anionic ligand binding and changes the catalyst E₁/₂. These results show how combining scaling relationships is a powerful way toward improved electrocatalysis.

INTRODUCTION

Improving the rates and efficiencies of electrocatalytic reactions is critical to the development of chemical-to-electrical energy conversion technologies. The oxygen reduction reaction (ORR)—the combination of dioxygen (O₂), protons, and electrons to give water—is an important example common to fuel cells (1, 2). Because current ORR technologies use platinum catalysts, current research seeks replacements sourced from earth-abundant materials (3). One approach to developing catalytic systems with high rates (turnover frequencies (TOFs)) and high efficiencies (low overpotentials, η) has been the development of scaling relationships. The properties of heterogeneous electrocatalysts can often be understood and even predicted using a single scaling descriptor, typically the energy of substrate binding to the catalyst surface (4). In contrast, soluble molecular electrocatalysts, which can be studied in more atomistic and mechanistic detail, follow multiple kinetic/thermodynamic scaling relationships (5). These describe how the maximum TOFs [log(TOFmax)] scale with the various terms contributing to the effective overpotential (ηeff) (5–8) (see the Supplementary Materials). To date, the reported log(TOFmax)/ηeff relationships always describe a trade-off: Faster catalysis is only achieved at lower efficiencies (higher ηeff).

We report here an inverse scaling relationship, one that allows for faster rates at higher efficiencies, using a polycationic iron porphyrin catalyst with buffered weak acids in acetonitrile (MeCN) (Fig. 1A). With acetic acid/acetate buffer (AcOH/AcO⁻), this system achieves a TOFmax of 170 s⁻¹ at 0.54 V ηeff, which is ~10⁴ faster than any previously reported molecular ORR catalyst at this ηeff (2, 9). As described below, this unprecedented result is predictable by combining two kinetic/thermodynamic scaling relationships. The coupling of the two scaling relationships is a result of the electrostatic ligand design of 1 that enables cooperativity between the catalyst and buffer.

RESULTS

Iron αβββ-tetra(o-N,N’,N”-trimethylanilinium)porphyrin (1) was prepared as reported (10) and was isolated and characterized as the di-aquo, penta-triflate salt of the αβββ isomer by single-crystal x-ray diffraction (Fig. 1C). Cyclic voltammograms (CVs) of 1 in MeCN containing 0.1 M tetra-n-butylammonium tetrafluoroborate electrolyte (n-Bu₄N][BF₄]) showed three reversible redox features (Fe³⁺/²⁺, Fe²⁺/¹⁺, and Fe¹⁺/⁰) (10). Solutions containing 1, O₂, and buffered acid (1:1 [HA]/[A⁻]) showed a large, irreversible current at the Fe³⁺/²⁺ reduction potential, indicative of catalysis (5, 7, 11). All CV experiments were buffered (1:1 acid-to-conjugate base) to define the ORR equilibrium potential and ηeff, which was calculated using Eq. 1 (5, 7, 11).

Measurements of the H₂O/H₂O₂ selectivity were performed using rotating ring disk electrochemistry (RRDE). In all cases, electrocatalysis was found to give <20% H₂O₂ (see the Supplementary Materials). This high selectivity for H₂O is similar to other iron porphyrin catalysts under comparable conditions (5, 7, 11). With AcOH/AcO⁻ buffer, catalysis occurs at potentials below the equilibrium potential for O₂ to H₂O and thus requires thermodynamic selectivity for H₂O [cf., (9)].

TOFmax values were determined from the catalytic CVs by foot-of-the-wave analysis (FOWA), Eq. 2 [Table 1; (5, 11, 12)]. This widely used approach normalizes the catalytic currents (iₖ) to the noncatalytic peak current of the Fe³⁺/²⁺ couple (iₚ) at potentials (E) near the onset of the catalytic wave, where complications such as substrate depletion are minimized. Given the high selectivity for H₂O, ncat was taken to be 4 (electrons per O₂ reduced), and the conservative σ = 1 value was used (see the Supplementary Materials). Measuring TOFmax values under different conditions showed that the catalytic rate law was first order in [1], P₂O₂, and [HA], similar to other Fe(por) catalysts (5, 7, 11). [HA] is the concentration of free acid after considering homoconjugation (see the Supplementary Materials). These results implicate a catalytic mechanism of (i) initial reduction of [Fe³⁺(por)]⁰ to Fe²⁺(por), (ii) pre-equilibrium O₂ binding to form the superoxo complex, Fe³⁺(por) (O₂•⁻), and (iii) rate-limiting proton transfer to form [Fe³⁺(por) (O₂H⁺)]⁺ (Fig. 1B). Experimental and computational studies of this mechanism are reported in (5, 11, 12).

ηeff = E⁰ ORR − E₁/₂ − 2.303RT ncat F log ηeff = − (0.0592V) p Kcat

(1)

iₖ iₚ = 2.24 ncat² E⁰ TOFmax

RT

1 + exp RT (E − E₁/₂)

(2)
electrocatalysis (Fig. 1A, blue dashed line), a previously unrealized goal in molecular
K\(_a\) = 23.5) gave markedly improved catalysis: a faster
(AcOH, p\(_\text{addition} \) (Fig. 1A, black diamond) (Fig. 1A). In contrast, using acetic acid
previously reported for Fe(por) electrocatalysts under similar con-
tricts previously derived log(TOF max)/\( \eta_{\text{eff}} \) space, which
\( \eta_{\text{eff}}, \log(\text{TOF max}) \) data point roughly fits the scaling relationships
is unique among the Fe(por) series because changing
(Fe, orange; N, blue; C, white; O, red; H atoms omitted for clarity; thermal ellipsoids at
50% probability).

Table 1. Properties of catalytic systems with 1 and different buffers. Experimental conditions: 0.1 mM 1, 0.1 M buffer (1:1 HA/A or HB+/B), 0.1 M [n-Bu4N][BF4] in MeCN (~15 mM H2O), 1 atm O2, [DMF-H+]N′-dimethylformamidium triflate; TFAH, trifluoroacetic acid; [Lut-H+]lutidinium tetrafluoroborate; SalOH, salicylic acid; BzOH, benzoic acid; AcOH, acetic acid.

| Buffer          | pK\(_a\)* | \( E_{1/2}(\text{Fe}^{III}/\text{II}) \)† | \( \eta_{\text{eff}} \)§ | TOF max (s\(^{-1}\))‡ | log(TOF max) |
|-----------------|----------|--------------------------------------|-------------------|------------------|--------------|
| None            | –        | –0.295                               | –                 | –                | –            |
| [DMF-H+]/DMF    | 6.1      | –0.25*                               | 1.16              | 8.5              | 0.91         |
| TFAH/TFA\(^-\)  | 12.6     | –0.349                               | 0.88              | 3.2              | 0.51         |
| [Lut-H+]/Lut    | 14.1     | –0.23*                               | 0.68              | 0.07             | –1.17        |
| SalOH/SalO\(^-\)| 16.7     | –0.536                               | 0.82              | 12               | 1.08         |
| BzOH/BzO\(^-\)  | 21.5     | –0.653                               | 0.67              | 63               | 1.80         |
| AcOH/AcO\(^-\)  | 23.5     | –0.651                               | 0.54              | 170              | 2.23         |

*See the Supplementary Materials. †\( E_{1/2}(\text{Fe}^{III}/\text{II}) \) reduction potential (versus ferrocenium/ferrocene, under Ar, with 100 mM buffer). ‡From Eq. 2; ±0.02 V.

ORR catalysis was studied in MeCN using 1 and a series of buffers. Using N,N’-dimethylformamidinium triflate ([DMF-H\(^+\)]OTf; pK\(_a\) = 6.1, where K\(_a\) is the acid dissociation constant), TOF max = 8.5 s\(^{-1}\) and \( \eta_{\text{eff}} = 1.16 \) V (Table 1). Within the log(TOF\(_\text{max}\))/\( \eta_{\text{eff}} \) space, this
\( \eta_{\text{eff}}, \log(\text{TOF max}) \) data point roughly fits the scaling relationships
previously reported for Fe(por) electrocatalysts under similar conditions (Fig. 1A, black diamond) (5). In contrast, using acetic acid (AcOH, pK\(_a\) = 23.5) gave markedly improved catalysis: a faster
TOF max (170 s\(^{-1}\) at less than the \( \eta_{\text{eff}} \) (0.54 V). This result con-
trads previously derived log(TOF\(_\text{max}\))/\( \eta_{\text{eff}} \) relationships, which
always predict that a lower overpotential will give a slower rate, as
seen for iron tetra-arylporphyrins (see the Supplementary Materials
(7). Catalysis with 1 and trifluoroacetic acid (TFAH), salicylic acid
(SalOH), or benzoic acid (BzOH) similarly show improvements in
both TOF\(_\text{max}\) and \( \eta_{\text{eff}} \) versus the [DMF-H\(^+\)] point (Fig. 1A, blue dia-
monds, and Table 1). Together, the carboxylic acid \( \eta_{\text{eff}}, \log(\text{TOF max}) \)
points define an unexpected inverse log(TOF\(_\text{max}\))/\( \eta_{\text{eff}} \) relationship (Fig. 1A, blue dashed line), a previously unrealized goal in molecular
electrocatalysis (5, 13, 14). The following sections describe a model
that explains this inverse relationship as the sum of two known
kinetic/thermodynamic scaling relationships (7).

DISCUSSION
Catalyst 1 is unique among the Fe(por) series because changing
the buffer notably affects two properties of the catalytic system:
the acid pK\(_a\) and also the catalyst \( E_{1/2} \), due to carboxylate bind-
ing. Since both pK\(_a\) and \( E_{1/2} \) each modulate the TOF max and \( \eta_{\text{eff}} \)
in different ways [see the Supplementary Materials and (5, 7)
for further discussion], a single scaling relationship is not ade-
quate for predicting the composite changes in log(OF max)/
\( \eta_{\text{eff}} \). Instead, the scaling relationships—which only establish
directionality within the log(OF max)/\( \eta_{\text{eff}} \) space—must be recast as vectors. Vectors have directionality and length and are
additive.

The Nernst equation for this ORR reaction (Eq. 1) shows that
a one-unit increase in acid pK\(_a\) decreases \( \eta_{\text{eff}} \) by 0.059 V: \( \Delta \eta_{\text{eff}} = -(0.059 \text{ V}) \Delta \text{pK}_a \) (15, 16). In iron porphyrin ORR catalysis, proton-
ation is the rate-determining step, and the TOF max varies with pK\(_a\) according to the Brønsted equation: \( \Delta \log(\text{TOF max}) = -\alpha(\Delta \text{pK}_a) \), where \( \alpha \approx 0.3 \) (5, 7, 11). For the range of carboxylate buffers used here,
TFAH/TFA\(^-\) to AcOH/AcO\(^-\), the change in pK\(_a\) is 10.9. Therefore,
with \( \alpha = 0.3 \) and holding all other properties constant, the changes in
\( \eta_{\text{eff}} \) and log(OF max) associated with the 10.9 pK\(_a\) shift can be
described as the vector $\mathbf{v}_{pK_a}$ in the $[\eta_{eff}, \log(\text{TOF}_{\text{max}})]$ space (Eq. 3, dec stands for decade in $\text{TOF}_{\text{max}}$; Fig. 3).

$$\mathbf{v}_{pK_a} = \langle \Delta \eta_{eff}, \Delta \log(\text{TOF}_{\text{max}}) \rangle = \langle -0.059(\Delta pK_a), -\alpha(\Delta pK_a) \rangle \approx \langle -0.64 \text{ V}, -3.3 \text{ dec} \rangle$$ (3)

In addition to acid $pK_a$, the buffer identity also affects the $E_{1/2}$. When buffered carboxylic acids are titrated into solutions of 1, the catalyst $E_{1/2}$ shifts negatively, by as much as 350 mV (Fig. 2A). This shift results from the conjugate base of the buffer (e.g., acetate) binding more strongly to the Fe(III) versus the Fe(II) form of 1. This behavior was documented and explained many years ago for chloride binding (17). At 0.1 M carboxylate buffer, the $E_{1/2}$ values vary linearly with acid $pK_a$ ($-28 \pm 1 \text{ mV/pK}_a$; Fig. 2B).

In contrast, buffers with cationic acids and neutral conjugate bases ([DMF-H$^+$]/DMF and [Lut-H$^+$]/Lut) give only small shifts in $E_{1/2}$ (<50 mV; Fig. 2B). This unique distinction is due to the highly cationic nature of 1, which enhances binding of anionic conjugate bases but not neutral ones (see the Supplementary Materials). Further supporting the importance of the cationic ligand, no notable change in catalyst $E_{1/2}$ was reported when iron tetrathiaporphyrin—an analog without the cationic trimethylanilinium groups—was combined with similar buffers under similar conditions (7).

Changes in $E_{1/2}$ also affect $\eta_{eff}$: $\Delta \eta_{eff} = -\Delta E_{1/2}$ (Eq. 2) (5, 7, 18). For the Fe(por) series, $\log(\text{TOF}_{\text{max}})$ has empirically been shown to change according to $\log(\text{TOF}_{\text{max}}) = -18.5 \text{ decade}/E_{1/2} (\text{V})$ (see the Supplementary Materials for further discussion) (7). Conceptually, this is because catalysts with more negative $E_{1/2}$ (and thus higher $\eta_{eff}$) bind O$_2$ more strongly and form more basic superoxide complexes, both of which lead to higher $\text{TOF}_{\text{max}}$ (5). As above, a vector $v_{E_{1/2}}$ can be used to represent the log(10)/$E_{1/2}$ scaling relation and the 0.302 V change in $E_{1/2}$ upon replacing TFAH/TFA$^-$ with AcOH/AcO$^-$ (Eq. 4 and Fig. 3).

$$\mathbf{v}_{E_{1/2}} = \langle \Delta \eta_{eff}, \Delta \log(\text{TOF}_{\text{max}}) \rangle = \langle -\Delta E_{1/2}, -18.5(\Delta E_{1/2}) \rangle$$ (4)

$$= \langle 0.30 \text{ V}, 5.6 \text{ dec} \rangle$$

Because the buffer affects both the acid $pK_a$ and catalyst $E_{1/2}$, both $v_{pK_a}$ and $v_{E_{1/2}}$ are needed to describe the changes in $\eta_{eff}$ and log(10)/$E_{1/2}$ (5, 7, 18). The effects of these changes are additive, described by the vector sum: $\mathbf{v}_{\text{sum}} = v_{pK_a} + v_{E_{1/2}}$ (Eq 5 and Fig. 3). While neither $v_{pK_a}$ nor $v_{E_{1/2}}$ alone fits the observed changes in log(10)/$E_{1/2}$, $v_{\text{sum}}$ predicts both the directionality and distance from the experimental TFAH coordinate to the AcOH coordinate—within a factor of 3 in $\text{TOF}_{\text{max}}$.

$$\mathbf{v}_{\text{sum}} = \mathbf{v}_{pK_a} + \mathbf{v}_{E_{1/2}} = \langle -0.34 \text{ V}, 2.3 \text{ dec} \rangle$$ (5)

Qualitatively, using a less acidic buffer improves the overall catalysis because of two factors. The change in $E_{1/2}$ causes a large gain in $\text{TOF}_{\text{max}}$ at relatively little cost to $\eta_{eff}$ (a steep scaling slope). In contrast, the concomitant changes in $pK_a$ decrease $\eta_{eff}$ substantially with only small losses in $\text{TOF}_{\text{max}}$ (a shallow slope). The combination is a faster $\text{TOF}_{\text{max}}$ at lower $\eta_{eff}$.

While $\mathbf{v}_{\text{sum}}$ is required to analyze the carboxylate buffer data, only $v_{pK_a}$ is needed to predict the change in log(10)/$E_{1/2}$ for the cationic buffers (e.g., [DMF-H$^+$]/DMF and [Lut-H$^+$]/Lut) (Fig. 3B). This is because the change in cationic buffers affects primarily the $pK_a$, while $\eta_{eff}$ is little affected on the $E_{1/2}$. The lack of change in $E_{1/2}$ highlights the key role of the cationic macrocycle, which enhances anion binding but not binding of a neutral ligand. Unlike CO$_2$ electroreduction by catalyst 1 (10), the remarkable log(10)/$E_{1/2}$ data reported here are not just a feature of the catalyst but rather the combination of the catalyst and buffer.

The summative or “tandem” scaling relationship approach developed here is unexpectedly accurate given that the contributing scaling relationships were obtained for a somewhat different set of catalysts, iron porphyrins with different meso-aryl substituents. Yet,
these scaling relationships quantitatively hold for the different axial ligands in the 1 + buffer systems examined here. Paradoxically, the tandem approach circumvents the limitations implied by prior molecular scaling relationships—that there is always a trade-off between $\text{TOF}_{\text{max}}$ and $\eta_{\text{eff}}$—because of the robustness and generality of the same “limiting” relationships.

Here, we show that kinetic/thermodynamic scaling relationships for molecular electrocatalysts can be additive and that this tandem scaling approach is a powerful and predictive way to improve multistep electrocatalytic processes. Specifically, summing multiple known scaling relationships predicted an unprecedented, inverse relationship that allowed for simultaneous improvements in both catalysis rates and efficiencies. This approach has yielded the best reported combination of $\text{TOF}_{\text{max}}$ and $\eta_{\text{eff}}$ for a soluble ORR catalyst.

The ability to combine scaling relationships should be applicable beyond this case study. Optimization by this method, and perhaps even inverse scaling, only requires that the catalytic reaction have at least two properties that affect the $\text{TOF}_{\text{max}}$ and $\eta_{\text{eff}}$ in different ways (different slopes). Most molecular electrocatalysis meets this two-descriptor requirement because their mechanisms have pre-equilibrium and rate-limiting steps with different stoichiometries and/or different relationships between the $k$, $K_{eq}$, and the $\Delta G^\circ$ for each step. ORR catalysis by 1 is a special case of the two-descriptor requirement because a single change to the system—changing the buffer—affects both the catalyst $E_{1/2}$ and the acid $pK_a$, each of which independently affects the pre-equilibrium and rate-determining steps. Ongoing work in our lab is extending this tandem scaling approach to simultaneous but independent changes in $E_{1/2}$ and $pK_a$, by changing both the catalyst and the buffer. A similar approach may also be applicable to heterogeneous electrocatalysis, since there is increasing recognition that those mechanisms may also have pre-equilibria and kinetic steps with different properties (19). Therefore, this method of combining scaling relationships may have ramifications for the development of both homogeneous and heterogeneous electrocatalysis of energy-important processes.

**MATERIALS AND METHODS**

**Instrumentation**

High-resolution mass spectrometry was performed using a Waters Xevo G2-XS QTof mass spectrometer. Ultraviolet-visible (UV-vis) optical spectra were recorded on an Agilent 8452 diode-array spectrometer and were collected using 1-cm path length cuvettes. Infrared (IR) spectrum was recorded on a Bruker Alpha Fourier transform infrared spectrophotometer equipped with an attenuated total reflectance (ATR) attachment. The electrochemical setup is described below. Information about the x-ray diffractometer is described below.

**Materials**

2-Nitrobenzaldehyde (Sigma-Aldrich, >95%), tin(II) chloride dihydrate (Sigma-Aldrich, >99%), aqueous hydrochloric acid [Macron, 36 to 38 weight % (wt %)], iron (II) bromide (Sigma-Aldrich, >98%), 2,6-lutidine (Sigma-Aldrich, ReagentPlus, >98%), formaldehyde (Sigma-Aldrich, 37 wt %, containing 10 to 15% methanol as stabilizer), sodium cyanoborohydride (Acros, >98%), methyltrifluoromethylsulfonate (MeOTf; Sigma-Aldrich, >98%), trifluoromethylsulfonic acid (Acros, 99%), neutral aluminum oxide (Sigma-Aldrich, >99%), silica (Sigma-Aldrich, >99%), sodium chloride (Sigma-Aldrich, >99%), sodium bicarbonate (Sigma-Aldrich, >99%), ammonium hydroxide solution (Sigma-Aldrich, 25 wt %), AcOH (Sigma-Aldrich, >99.9%), BzOH (Sigma-Aldrich, >99.5%), SalOH (Sigma-Aldrich, >99%), TFA (Sigma-Aldrich, ReagentPlus, >99%), tetra-n-butyllammonium acetate (Sigma-Aldrich, 97%), tetra-n-butyllammonium benzoate (Sigma-Aldrich, >99%), tetra-n-butyllammonium salicylate [Tokyo Chemical Industry (TCI), 98%], and sodium trifluoracetate (Sigma-Aldrich, 98%) were all used as received. All of the solid chemicals were stored in a N$_2$ glovebox when not in use.

Tetrahydrofuran, N,N$'$-dimethylformamide (DMF), MeCN, and diethyl ether were all degassed with argon and dried using a Pure Process Technology solvent system before use. Tetra-n-butyllammonium tetrafluoroborate (Acros, >98%) was stored in a desiccator containing Drierite (calcium sulfate). Bis(cyclopentadienyl)iron(II) (ferrocene or Fe; Sigma-Aldrich, 95%) was recrystallized two times from hexanes before being dried in a N$_2$ glovebox. Pyrrole (Acros, 99%) was freshly distilled immediately before each use. The Pur-A-Lyzer Mega Dialysis Kit [Sigma-Aldrich, 1 kDa molecular weight cut-off (MWCO)] was soaked in Milli-Q water for 1 hour before dialysis of 1. Dioxygen (Airgas, Ultra High Purity) and argon (Airgas, Ultra High Purity) were used as received. N,N$'$-dimethylformamidum triflate ([DMF-H]OTf) and 2,6-lutidinium triflate ([Lut-H]OTf) were synthesized using previously reported methods (20, 21).

**Synthesis of iron αβαβ-tetra(o-N,N',N'-trimethylanilinilinil) porphyrin (1)**

The synthesis of the known iron(III) αβαβ,β-5,10,15,20-tetra(ortho-N,N',N'-trimethylanilinilinil)-porphyrin penta(trifluoromethanesulfonate), 1, was slightly adapted from literature preparation (10). The αβαβ,β-5,10,15,20-tetra(ortho-aminoethyl)porphyrin hydrolysis was isolated chromatographically using the conditions described in (10). Later in the synthesis, during the final methylation reaction of iron(III) αβαβ,αβ,β-5,10,15,20-tetra(ortho-N,N',N'-dimethylanilinilinil)porphyrin chloride [chart S1, product 4, from reference (10)] to the fully quaternized product, 300-equivalent MeOTf were used in place of the reported 100 equivalents. Full quaternization was supported by the high-resolution mass spectrum (HRMS) which did not reveal any partially quaternized products and by cyclic voltammetry (see the Supplementary Materials). The product, which was collected in identical yields to the initial report, was purified by dialysis against Milli-Q water (10 ml of sample inside dialyzing bag, 250 ml of surrounding water replaced every 90 min for a total of 12 hours). The product was then recrystallized via slow vapor diffusion of diethyl ether into MeCN containing the iron porphyrin. This recrystallization method yielded crystals suitable for x-ray diffraction (see the Supplementary Materials). HRMS (electron spray ionization (ESI)/quadrupole orthogonal acceleration–time-of-flight) mass/charge ratio $(m/z)$: [(M$^+$ + 4OTf$^-$) calculated (calcd) for C$_{23}$H$_{16}$FeN$_6$(CF$_3$SO$_2$)$_3$ 1496.2373; found 1496.2493. [(M$^+$ + 3OTf$^-$) calcd for C$_{23}$H$_{16}$FeN$_6$(CF$_3$SO$_2$)$_3$ 767.6426; found 767.6395. UV-vis and IR spectra also agree with what was reported in reference (10), see the Supplementary Materials.

**Electrochemical methods**

CV was performed on a CH Instruments model 650D potentiostat. RRDE was conducted using a BASI Epsilon potentiostat and a Pine Instruments rotator (see the Supplementary Materials for further details). CVs used a 3-mm glassy carbon working electrode, a platinum wire counter electrode, and an Ag wire pseudoreference. The
Ag wire pseudoreference was prepared using the method reported by Dempsey et al. (22), where a silver wire was sanded and fit snugly into a capillary containing MeCN and 0.1 M [n-Bu₄N][BF₄]. The solution inside the capillary was separated from bulk solutions using a Vycor tip that had been mechanically sealed to the capillary using heat shrink tubing. The capillary was stored in a solution containing 0.1 M [n-Bu₄N][BF₄] in-between experiments. For analysis, each voltammogram was internally referenced to ferrocene. The glassy carbon working electrodes were polished after every voltammogram by vigorously polishing on a Buehler felt pad to a mirror-like finish using an alumina slurry (wetted 0.05-μm alumina powder). After polishing, the electrode was rinsed with water and MeCN. Internal resistance (IR) compensation was performed before each voltammogram using the CH Instruments integrated software. The typical resistance value was <60 ohm. Considering the typical amount of current passed during catalysis (~200 μA), the IR compensation shifted the potential response of <10 mV for most experiments.

All of the buffered electrochemical experiments were performed with a one-to-one buffer of acid and conjugate base. Throughout, we identify the initial buffer concentrations rather than expressly list the acid and conjugate base concentrations. For example, a “10 mM AcOH buffer” means that the solution was prepared to initially contain 10 mM AcOH and 10 mM AcO⁻. These values do not necessarily reflect the concentrations of nonhomoconjugated acid and base (see the Supplementary Materials).

Electrocatalytic, aerobic measurements were performed after sparging the buffered solution and headspace with O₂. The sparging was performed with pure O₂ at 1 atm unless otherwise noted. To minimize solvent evaporation and accompanying temperature changes, the O₂ was first sparged through an initial bubbler containing pure MeCN. Between each catalytic voltammogram, the working electrode was vigorously polished (see above). To ensure reproducibility, each voltammogram was also performed in duplicate, and the rates were averaged (see the Supplementary Materials). Unless otherwise specified, every voltammogram reported was collected at 0.1 V/s in a MeCN solution containing 0.1 mM 1 and 0.1 M [n-Bu₄N][BF₄].

**Section: Single-crystal x-ray diffraction**

Crystals of [Fe-ο-TMA]OTf₂•2H₂O (identification code 007b-17062) were grown by slow vapor diffusion of diethyl ether into MeCN on the benchtop. Two water molecules were identified as axial ligands in the solid-state structure. Low-temperature diffraction data (ω-scans) were collected on a Rigaku MicroMax-007HF diffractometer coupled to a Saturn994+ charge-coupled device detector with Cu Kα (λ = 1.54178 Å) for the structure of [Fe-ο-TMA]OTf₂•2H₂O. The diffraction images were processed and scaled using Rigaku Oxford Diffraction software. The structure was solved with SHELXT and was refined against F² on all data by full-matrix least squares with SHELXL (23). The data were refined as an inversion twin. The fractional volume contributions of the second twin component were refined to a value of 0.432(18). All nonhydrogen atoms were refined anisotropically. Hydrogen atoms were included in the model at geometrically calculated positions and refined using a riding model. The isotropic displacement parameters of all hydrogen atoms were fixed 1.2 times the U value of the atoms to which they are linked (1.5 times for methyl groups). The Fe-O distance was measured to be 2.101(8) Å, which suggested the assignment as water.

The asymmetric unit of the crystallographic model contains 0.25 of the formula unit C₅₆H₆₄FeN₈O₂•5(CF₃O₃S). The iron resides on the crystallographic 4 rotation axis at (0, ½, ¼). The asymmetric unit also contains a quarter of the porphyrin and 1.25 triflates. One triflate site is on a general position (x,y,z), and the other site is near the crystallographic 4 rotation axis (½, ½, ½).

The triflate on the general position is disordered over two sites. The thermal ellipsoids for the triflate atoms were restrained to behave as rigid bodies. The site occupancies were fixed at 0.50, and all chemically equivalent 1,2 and 1,3 distances were restrained to be similar. The other triflate was disordered with respect to the 4 rotation axis. This modeled triflate was placed as a constrained, rigid group based on the difference map (see the Supplementary Materials). The special position constraints were suppressed, and the model occupancy was fixed at 0.25. The program SQUEEZE (24) was used to compensate for the contribution of disordered solvents contained in voids within the crystal lattice from the diffraction intensities. This procedure was applied to the data file, and the submitted model is based on the solvent removed data. On the basis of the total electron density found in the voids (181 e/Å³), it is possible that some combination of crystallization solvents is present in the unit cell. See “_platon_squeeze_details” in this .cif for more information.

The full details of the x-ray structure determination (CIF) can be found in the Cambridge Crystallographic Data Center (CCDC), number 1947226 ([Fe-ο-TMA]OTf₂•2H₂O). These data can be obtained free of charge from the CCDC via www.ccdc.cam.ac.uk/data_request/cif.

**SUPPLEMENTARY MATERIALS**

Supplemental material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/11/eaaz3318/DC1

Section S1: Spectroscopic characterization of 1
Section S2: Voltammetry of 1
Section S3: UV-vis spectroscopy of 1 + varying buffers
Section S4: Homocoujugation
Section S5: Kinetic analysis
Section S6: Effective overpotential determination
Section S7: Selectivity for H₂O versus H₂O₂
Section S8: Conceptual background for E₁/₂ and pKₐ scaling relationships
Section S9: Single crystal x-ray structure
Fig. S1. Full high-resolution ESI mass spectrum of 1 with identified peaks, as labeled.
Fig. S2. High-resolution ESI mass spectrum and isotopic fits for the [M⁺ + 30OTF⁻] ion.
Fig. S3. UV-vis spectrum of 1 in N,N-dimethylformamide.
Fig. S4. IR spectrum of 1.
Fig. S5. A CV of an Ar-sparged solution of 1.
Fig. S6. Scan rate investigation of the Fe³⁺/Fe²⁺ redox couple of 1 in unbuffered solution.
Fig. S7. Voltammograms of an Ar-sparged solution of 1 before and after addition of buffered AcOH.
Fig. S8. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of AcOH buffer.
Fig. S9. Scan rate investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 in AcOH-buffered solution.
Fig. S10. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of BuOH buffer.
Fig. S11. Scan rate investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 in BuOH-buffered solution.
Fig. S12. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of SalOH buffer.
Fig. S13. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of SalOH-buffered solution.
Fig. S14. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of TFA buffer.
Fig. S15. Scan rate investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 in TFA-buffered solution.
Fig. S16. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of DMF-HDOTT buffer.
Fig. S17. Investigation of the Fe⁵⁺/Fe⁴⁺ redox couple of 1 with titrations of DMF-HDOTT-buffered solution.
Fig. S18. Changes in E₁/₂(Fe³⁺/Fe²⁺) with varying buffers (and concentrations).
Fig. S19. CVs of an O₂-sparged MeCN solution containing 100 mM [AcOH] buffer before (black) and after (blue) adding 30 μM 1.
Fig. S20. Voltammograms of 1 with AcOH buffer under various solution conditions.
Fig. S21. Voltammograms of 1 with BuOH buffer under various solution conditions.
Fig. S22. Voltammograms of 1 with SalOH buffer under various solution conditions.
Fig. S23. Voltammograms of 1 with TFA buffer under various solution conditions.
Fig. S24. Voltammograms of 1 with DMF-HDOT buffer under various solution conditions.
Fig. S25. Voltammograms of 1 with DMF-HDOTT buffer under various solution conditions.
Fig. S26. Rinse tests for all of the buffers used in this study.
max values for ~0.05 mM reported in this study; errors are ±0.005 V.

-TMA]OTf 5 •2H2O.

and AcOH buffer.

Fig. S37. RRDE analysis for the ORR catalyzed by and BzOH buffer.

1

Fig. S36. RRDE analysis for the ORR catalyzed by using SalOH buffer.

1

Fig. S35. RRDE analysis for the ORR catalyzed by 1 using SAOH buffer.

1

Fig. S34. RRDE analysis for the ORR catalyzed by 1 using TFAH buffer.

1

Fig. S33. RRDE analysis for the ORR catalyzed by using [DMF-H]OTf buffer.

Fig. S32. RRDE analysis using ferrocene to estimate collection efficiencies.

Table S11. Torsion angles (°) for [Fe-o-TMA]OTf 5 •2H2O.

Table S10. Hydrogen coordinates (×10⁴) and isotropic displacement parameters (Å² × 10³) for [Fe-o-TMA]OTf 5 •2H2O.

Table S9. Anisotropic displacement parameters (Å² × 10³) for [Fe-o-TMA]OTf 5 •2H2O.

Table S8. Bond lengths (Å) and angles (°) for [Fe-o-TMA]OTf 5 •2H2O.

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