Electrocatalytic Ammonia Oxidation Mediated by a Polypyridyl Iron Catalyst

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3.1 Introduction

Ammonia is produced on a massive scale globally by industrial nitrogen fixation, primarily for use as fertilizer. New ammonia synthesis technologies may yet enable other vectors for ammonia use, for example as a transportation fuel. Accordingly, there are substantial efforts underway to explore whether electrocatalytic N₂-to-NH₃ (N₂RR) conversion, potentially coupled with renewable resources that generate electricity, could provide a new ammonia synthesis pathway (Equation 3.1) to be exploited in renewably formed fuel storage, transport, and on-site use.¹

\[
\text{N}_2 + 6 \text{H}^+ + 6 \text{e}^- \rightleftharpoons 2 \text{NH}_3 \quad \text{(Equation 3.1)}
\]

Commensurate with these goals, there has been long-standing interest in exploring heterogeneous catalysts for selective ammonia oxidation (AO), the microscopic reverse of N₂-to-NH₃ conversion (Equation 3.1).² The realization of selective electrocatalysts for AO in principle could enable fuel cell applications for ammonia, akin to those currently being practiced or explored for hydrogen and methanol.³

Homogeneous (electro)catalysts for N₂RR and AO have much to offer in terms of fundamental mechanistic studies, and possibly longer-term practical applications if robust hybrid catalyst/electrode architectures can be realized. Indeed, the field of N₂RR catalysis mediated by molecular systems has witnessed a surge of activity in the past 10–15 years, in part motivated by a desire to explore hypotheses germane to the inorganic mechanism of biological nitrogen fixation.⁴ This has been matched by significant progress in defining catalysts (e.g., featuring Mo or Fe) that operate with (comparatively) impressive turnovers.
and selectivities, and in fundamental mechanistic understanding of how these transformations occur. Interest in translating this progress to electrocatalytic N₂RR using synthetic coordination complexes is growing.

By contrast, homogeneous approaches to ammonia oxidation have been slower to attract significant attention from the coordination chemistry community. This is surprising given that biological ammonia oxidation, mediated by both aerobic and anaerobic ammonia-oxidizing bacteria, involves steps thought to occur at iron, copper, or molybdenum active sites, and is essential to the global nitrogen cycle. As a six-electron process with substantial kinetic barriers, this process presents fascinating challenges in energy conversion, electrochemistry, and coordination chemistry. Challenges associated with catalytically converting NH₃ to N₂ and proton/electron equivalents at a well-defined active site include the fact that NH₃ is a strong σ-donor ligand that features thermally robust N–H bonds (BDFE₇N–H = 99.4 kcal/mol).

Despite the comparatively limited activity in this area, stoichiometric oxidation of ammonia is well preceded. For instance, in 1979 Buhr and Taube initiated the study of ammonia oxidation by molecular complexes when they reported both chemical and electrochemical oxidation of ammonia in aqueous solution by [Os(NH₃)₅(CO)]²⁺ to form N₂ in the μ-N₂-bridged product [(Os(NH₃)₄(CO))₂N₂]⁴⁺ (Chart 3.1, A). Relatedly, Thompson and Meyer reported the stoichiometric electrochemical oxidation of ammonia to nitrite and nitrate by [(trpy)(bpy)Ru(NH₃)]²⁺ (trpy = 2,2′:6′,2′′-terpyridine, bpy = 2,2′-bipyridine) in aqueous solution (Chart 3.1, B), and reported electrochemical (though not electrocatalytic) ammonia oxidation to N₂ by a similar ruthenium polypyridyl system,
[(bpy)$_2$(NH$_3$)RuORu(NH$_3$)(bpy)$_2$]$^{2+}$. Collman and coworkers have studied chemical and electrochemical oxidation of ammonia at cofacial ruthenium porphyrins via hydrazine and diazene intermediates. Most recently, Hamann, Smith, and coworkers demonstrated that modification of the Meyer system provides a bona fide electrocatalyst for ammonia oxidation to N$_2$, reporting the liberation of 2.1 eq N$_2$ at 0.2 V vs Fc/Fc$^+$ (0.03 V onset potential) under the conditions used (Chart 3.1, C).

There has also been growing interest in exploring molecular catalysts for ammonia oxidation that operate via net H-atom abstraction chemistry. In this context, a number of fascinating recent reports describe H-atom abstraction (HAA) reactions from coordinated NH$_3$, which can lead to M(NH$_2$), M(NH), M(N), and in certain cases N$_2$ (via nitride coupling). Catalytic ammonia oxidation via HAA has also been recently demonstrated using a ruthenium complex and 2,4,6-tri(tert-butyl)phenoxy radical by Mock and coworkers.
Similarly, Nishibayashi and coworkers recently reported catalytic chemical oxidation of ammonia, again using a ruthenium system (Chart 3.1, E).16

Given the paucity of data available describing electrocatalytic ammonia oxidation, and the fact that biologically relevant first row metals such as iron have yet to be described for this process, we targeted the study of promising candidate iron (electro)catalysts as models well-suited to mechanistic interrogation. Noting that a number of polypyridyl iron complexes have been studied as potential water oxidation catalysts (WOC’s),17 we felt such systems might provide a prudent starting point.

In this context, we now describe that a previously reported iron complex featuring a tetradeutate polypyridyl supporting ligand (TPA) and two cis-labile sites, [(TPA)Fe(MeCN)2]2+,18 is highly active for electrocatalytic ammonia oxidation in acetonitrile under an applied bias. Important features of this system are that NH3 can reversibly bind at the two labile sites, NH3 itself is able to promote the proton-transfer steps needed for catalytic activity, and up to 16 eq of N2 (per Fe) have been generated from NH3 using this catalyst via controlled potential coulometry (CPC) experiments. At present, this is the highest TON to be verified for a molecular AO system. More importantly, the system is electrochemically well behaved and hence amenable to detailed study by a range of electrochemical experiments. Our data point to an observed rate constant, $k_{obs}$, for NH3 oxidation of $\sim 10^7$ M$^{-1}$s$^{-1}$, with the catalytic process starting at an onset potential of $\sim$0.7 V vs Fe/Fe$^+$, whose rate-determining step (RDS) is first order in [(TPA)Fe(L)2]$^{2+}$ and NH3. The available data also allow us to suggest [(TPA)Fe$^{III}$(NH2)(NH3)]$^{2+}$ and [(TPA)Fe$^{IV}$(NH2)(NH3)]$^{3+}$ species as early intermediates en route to N–N bond formation.
3.2 Results and Discussion

3.2.1 Electrocatalytic conversion of NH₃ to N₂ and H₂. The cyclic voltammogram of [(TPA)Fe(MeCN)₂]OTf₂ in acetonitrile using a boron-doped diamond (BDD) working electrode (see Figure 3.1A,B) has one reversible feature at 0.75 V vs Fc/Fc⁺ corresponding to the Fe²⁺/Fe³⁺ redox couple (Figure 3.1A). Two new redox features appeared (E₁ and E₂) in the presence of increasing amounts of added NH₃, and the reversible feature corresponding to [(TPA)Fe(MeCN)₂]OTf₂ disappeared. Moreover, the wave at E₂ continuously increased with increasing ammonia concentration. As a reference, when 100 equivalents of NH₃ were added to the [(TPA)Fe(MeCN)₂]OTf₂ solution in acetonitrile, E₁ and E₂ were centered at 0.4 V and 1.1 V, respectively, and E₂ featured dramatically increased current relative to that.

Figure 3.1. Electrochemical data. (A) Cyclic voltammograms in acetonitrile solutions of background AO on BDD in presence of 0.25 M NH₃ (black), 2.5 mM [(TPA)Fe(MeCN)₂]OTf₂ in the absence of NH₃ (red), and AO by [(TPA)Fe(MeCN)₂]OTf₂ with 0.25 M NH₃ (green). (B) Cyclic voltammograms in acetonitrile with 50 mM NH₄OTf and 0.5 mM [(TPA)Fe(MeCN)₂]OTf₂ with varying [NH₃]. The inset highlights the shift in E₁ with [NH₃]. CVs were recorded with a BDD disk electrode, a Pt counter electrode, and a custom Ag/AgOTf reference electrode corrected for the Fc/Fc⁺ couple.
observed for the one-electron Fe$^{2+}$/Fe$^{3+}$ redox couple. These observations are consistent with fast electrocatalytic ammonia oxidation at $E_2$.

To assess whether catalytic NH$_3$ oxidation occurs at the irreversible $E_2$ wave, CPC was performed using a BDD plate working electrode (1 cm$^2$) and a Pt counter electrode to facilitate H$_2$ evolution, and the headspace was analyzed via gas chromatography. CPC of a 0.5 mM [(TPA)Fe(MeCN)$_2$]OTf$_2$ acetonitrile solution with 65 mM NH$_3$ was performed at 1.1 V vs Fc/Fc$^+$. After 18 h, 33 C of charge had passed; headspace analysis indicated Faradaic efficiencies of 80% and 70% for the production of N$_2$ and H$_2$ (±10% error in gas quantification), respectively, correlating with the production of up to 16 equivalents of N$_2$. This turnover number is the highest yet reported for a molecular NH$_3$ oxidation (electro)catalyst. The measured H$_2$:N$_2$ ratio was 2.6:1, in good agreement with the 3:1 ratio expected for NH$_3$. When isotopically enriched $^{15}$NH$_4$OTf and $^{15}$NH$_3$ were employed, only $^{30}$N$_2$ was observed by GC-MS, confirming ammonia to be the source of detected N$_2$ (see SI for details).

Further CPC experiments in the absence of a (TPA)Fe source, at the same applied potential of 1.1 V vs Fc/Fc$^+$, suggest a critical role for (TPA)Fe-species in the catalysis. When CPC of a 65 mM NH$_3$ solution was performed, only 1.1 C of charge was passed, and only 0.5 eq N$_2$ were detected. Furthermore, when [(TPA)Fe(MeCN)$_2$]OTf$_2$ was replaced with FeOTf$_2$ as the precatalyst, only 1.9 C of charge were passed along with 1.0 eq N$_2$ being detected in the headspace.

3.2.2 Investigation of catalyst stability. Over multiple independent CPC experiments, the current after 18 h at an applied potential of 1.1 V vs Fc/Fc$^+$ had greatly attenuated. Such a
decrease in current typically suggests either catalyst decomposition or electrode passivation.\textsuperscript{20} To study this phenomenon, X-ray photoelectron spectroscopy was performed on the BDD working electrode after CPC to try to identify potential degradation products. During CPC with 65 mM NH\textsubscript{3} and no added iron precatalyst, the current dropped to 10\% of the initial current after only 3 h. XPS analysis of the BDD electrode revealed incorporation of nitrogen onto the electrode surface (see SI). This process passivates the electrode surface, as confirmed by measurements of the ferrocene/ferrocenium redox couple before and after CPC. This passivation is hence one process by which the catalysis can be arrested in the presence of NH\textsubscript{3} using BDD electrodes.

Although the chronoamperogram with [(TPA)Fe(MeCN)\textsubscript{2}]OTf\textsubscript{2} as a precatalyst in the presence of NH\textsubscript{3} also showed a time-dependent decrease in current, a much longer time scale, \textasciitilde15 h, was required to reach 10\% of the initial current. Measurement of the working electrode after CPC again revealed incorporation of nitrogen, but now also iron, on the surface. The detection of iron on the electrode surface by XPS raises the possibility that a heterogeneous iron catalyst may be responsible for (or contribute to) the electrocatalytic AO discussed above. Indeed, it is difficult, if not impossible, to reliably discount a contributing role for heterogenous (electro)catalysis.\textsuperscript{21} However, several lines of evidence lead us to suggest that a molecular, TPA-ligated iron catalyst dominates the aforementioned AO behavior. In the chronoamperogram for AO by the precatalyst [(TPA)Fe(MeCN)\textsubscript{2}]OTf\textsubscript{2}, no induction period was observed. Furthermore, a rinse test was performed with the BDD electrode following CPC using a fresh acetonitrile-ammonia solution. No catalytic current was passed in this case, ruling out a heterogeneous catalyst that is strongly attached to the electrode. Measurement of the catalyst solution after CPC with a clean BDD disk electrode
indicated a catalytic wave of similar intensity to that observed prior to CPC. To demonstrate the relative stability of the TPA-ligated Fe-species under catalytic conditions, 50 cycles of CV were performed, Figure 3.2A. Almost no reduction in catalytic current appeared between the first and last scan. While these experiments cannot reliably discount the possibility of a very rapidly formed, loosely bound and highly active heterogeneous catalyst forming under CPC experiments, we find such a scenario improbable.

![Graph](image)

**Figure 3.2.** Cycling of (A) 2.5 mM [(TPA)Fe(MeCN)₂]OTf₂, or (B) 2.5 mM FeOTf₂, with a BDD disk electrode in an acetonitrile solution containing 50 mM NH₃ and 50 mM NH₄OTf. Pt wire was used as the counter electrode with a custom Ag/AgOTf reference electrode.

To probe this issue further, [(TPA)Fe(MeCN)₂]OTf₂ was replaced by FeOTf₂ as the precatalyst during CPC with ammonia. In this case, the current rapidly dropped to 10% of the initial current after only 1 h. After CPC, the XPS spectrum of the BDD electrode was almost identical to the spectrum obtained after CPC with [(TPA)Fe(MeCN)₂]OTf₂. 50 cycles of CV were also performed with FeOTf₂ in ammonia, Figure 3.2B. A continuous decrease in peak current, correlated with passivation of the electrode, was observed. In acetonitrile-ammonia solutions, we presume FeOTf₂ exists primarily as [Fe(NH₃)₆]OTf₂. The decrease
in current observed by CPC and CV, in addition to the iron observed on the electrode surface by XPS, establish the instability of \([\text{Fe(NH}_3)_6\text{OTf}_2]\) under the present conditions.

Given the instability of \([\text{Fe(NH}_3)_6\text{OTf}_2]\) under electrocatalytic conditions, we wondered if the source of the iron observed on the electrode surface post-CPC could be due to partial demetallation of TPA-ligated Fe species by NH\(_3\), thereby forming \(\text{Fe(NH}_3)_6^{2+}\), which can then degrade at the BDD electrode. The speciation of \([\text{(TPA)Fe(L)}_2\text{OTf}_2]\) with varying ammonia concentrations was therefore studied by UV-vis spectroscopy (Figure 3.3A,B). A plot of absorbance at 400 nm vs \([\text{NH}_3]\), Figure 3.3A, displays three distinct regions of different slopes. We assign these as regions of equilibria corresponding to the successive displacement by NH\(_3\) of one MeCN (1–10 eq NH\(_3\)), a second MeCN (10–200 eq NH\(_3\)), and finally the additional displacement of TPA by NH\(_3\) to form \(\text{Fe(NH}_3)_6^{2+}\). In accord with these equilibria being reversible, the addition of free TPA to \(\text{FeOTf}_2\) in MeCN with

**Figure 3.3.** UV-vis monitoring of speciation. (A) Plot of absorbance at \(\lambda_{\text{max}} = 400\) nm versus \([\text{NH}_3]\) demonstrating regions corresponding to mono-ammine, bis-ammine, and hexakis-ammine iron complexes. (B) UV-vis spectrum of 0.2 mM \([\text{(TPA)Fe(MeCN)}_2\text{OTf}_2]\) with 0, 20, 100, or 1500 eq NH\(_3\) added in acetonitrile.
excess NH₃, to reproduce the electrocatalytic conditions, produces the \( E_1 \) and \( E_2 \) redox features as well as the UV-vis spectrum associated with \([\text{(TPA)Fe(NH}_3)_2]\)^{2+}; in the absence of FeOTf₂, an irreversible oxidation wave for free TPA is instead observed at ~ 1.0 V. Consistent with the viability of \([\text{(TPA)Fe(NH}_3)_2]\)OTf₂, we were able to obtain its solid-state X-ray structure via crystals grown by diffusing NH₃ gas into a THF solution of dissolved (TPA)FeOTf₂. The solid-state structure of \([\text{(TPA)Fe(NH}_3)_2]\)OTf₂ features four independent molecules in the asymmetric unit and interestingly, bond lengths that are consistent with the presence of both high- and low-spin iron centers (two of each type; see Figure 3.4 and SI). ²² Given that \([\text{(TPA)Fe(NH}_3)_2]\)OTf₂ is structurally well-defined, as is \([\text{(TPA)Fe(MeCN)}_2]\)OTf₂, ¹⁸ an intermediate structure, \([\text{(TPA)Fe(NH}_3)(\text{MeCN})]\)OTf₂, is highly plausible and we presume the primary species present in the 1–10 eq NH₃ region.

Although the aforementioned electrocatalytic conditions include 130 eq NH₃, a regime in which \([\text{(TPA)Fe(NH}_3)_2]\)OTf₂ dominates, there is likely a small population of \([\text{Fe(NH}_3)_6]\)^{2+} and also free TPA under these conditions. The source of iron on the electrode presumably results as \([\text{Fe(NH}_3)_6]\)^{2+} is continuously degraded, shifting the equilibrium toward further demetallation. Fortunately, degradation is slow, as evidenced by CV cycling experiments, the chronoamperogram as well as CVs and UV-vis acquired before and after CPC, (see SI).
3.2.3 Mechanistic insight into the first redox process ($E_1$). The sequential addition of NH$_3$ to [(TPA)Fe(L)$_2$]$^{2+}$ increased the catalytic current at $E_2$ ($E_{cat}$), Figure 3.1B. Another change observed in the cyclic voltammogram of [(TPA)Fe(L)$_2$]$^{2+}$ in MeCN with increasing amounts of ammonia is a shift in the potential of the first redox process, $E_1$, toward more cathodic values. This observation indicates that the electron transfer step (E) in the oxidation is
coupled to a chemical step (C) involving NH$_3$ as a reactant. This EC mechanism is further supported by an increasing irreversibility at $E_1$ concomitant with the appearance of a new reduction peak $E_1'$ at $-0.4$ V (Figure 3.5A), arising from the formation of a new species as [NH$_3$] is increased.

**Scheme 3.1.** Equilibria relevant to substitution of acetonitrile and TPA by ammonia in acetonitrile solution for [(TPA)Fe(L)$_2$]$^{2+}$.

![Scheme 3.1](image)

Interestingly, only one oxidation peak was observed at lower ammonia concentrations. This can be rationalized via a fast equilibrium between [(TPA)Fe(MeCN)(NH$_3$)]$^{2+}$ and [(TPA)Fe(NH$_3$)$_2$]$^{2+}$ as described in Scheme 3.1. The bis-amine complex is expected to have a more cathodically shifted oxidation potential (NH$_3$ being a stronger donor than MeCN). Thus, as the population of [(TPA)Fe(NH$_3$)$_2$]$^{2+}$ is oxidized to Fe(III) near the electrode, the equilibrium shifts, driving coordination of a second NH$_3$ eq to [(TPA)Fe(MeCN)(NH$_3$)]$^{2+}$; the latter species should still be primarily in the Fe(II) oxidation state as it is expected to have a redox potential $\sim$150 mV more positive than the [(TPA)Fe(NH$_3$)$_2$]$^{2+}$. Rapid solvent equilibration is expected for high spin iron(II), on the order of $\sim$10$^4$–10$^6$ s$^{-1}$.24
The voltammetric response of an EC mechanism is governed by the pseudo-first order equilibrium constant of the chemical step (K), as well as the competition between the pseudo-first order rate constant of the chemical step \( k' \) and diffusion.\(^{25,26} \) Under our present conditions, the first redox event \( E_1 \) shows quasi-reversible character typical for the KE regime in the kinetic zone diagram, where the redox potential is governed by the following equation:

\[
E_{ox} = E_{ox}^0 - \frac{RT}{F} \ln \left( \frac{k'_{1f}}{k'_{1b}} \right) \quad \text{(Equation 3.2)}
\]

\( F \) is the Faraday constant, \( R \) is the gas constant, \( T \) is the temperature, \( E_{ox} \) is the oxidation potential, \( E_{ox}^0 \) is the standard oxidation potential, and \( k'_{1f} \) and \( k'_{1b} \) are, respectively, the pseudo first order rate constants for the forward and backward chemical reaction. Using Equation 3.2, the change in peak potential with the natural logarithm of ammonia concentration reveals a linear dependence with a negative slope of \(-0.030\), in good agreement with the theoretical value of \(-0.026\) (Figure 3.5B). Due to its character as a Brønsted base, one plausible explanation for the role of NH\(_3\) in the chemical step following oxidation is the abstraction of a proton from one of the NH\(_3\) ligands coordinated to Fe(III). Such a mechanism would produce NH\(_4^+\) as a product, together with the oxidized iron complex \([(TPA)Fe^{III}(NH_3)(NH_2)]^{2+}\). Accordingly, the \( E_i \) potential with varying concentrations of NH\(_4^+\) at a fixed concentration of NH\(_3\) (Figure 3.5C) shows a linear relationship, with an observed slope of +0.024. This finding is in agreement with the predicted value of +0.026.
Hence, the proposed EC mechanism is consistent with the electrochemical behavior at $E_1$ as reactant, NH$_3$, and product, NH$_4^+$, concentrations are varied.

**Figure 3.5.** Evidence supporting an EC mechanism. (A) CV of an acetonitrile solution containing 0.5 mM [(TPA)Fe(MeCN)$_2$]$^{2+}$, 50 mM NH$_4$OTf, and 10 (black dashed trace) or 50 (solid green trace) eq NH$_3$. (B) Plot of the potential for the first redox event $E_1$ versus the natural logarithm of [NH$_3$]. (C) Plot of the potential for the first redox event $E_1$ versus the natural logarithm of [NH$_4^+$]. (D) CV of an acetonitrile solution containing 0.5 mM of [(TPA)Fe(MeCN)$_2$]$^{2+}$, 50 mM NH$_4$OTf, and 50 mM NH$_3$ at different scan rates. CVs recorded with a BDD disk electrode, a Pt counter electrode, and a custom Ag/AgOTf reference electrode.
Using the above logic, we assign the reduction peak $E_1'$, appearing at $-0.4$ V in Figure 3.5A, to the one-electron reduction of $[(TPA)Fe^{III}(NH_3)(NH_2)]^{2+}$. Upon its reduction, no return oxidation peak could be detected in the following anodic scan near $-0.4$ V, indicating fast protonation to regenerate $[(TPA)Fe^{II}(NH_3)_2]^{2+}$. This is consistent with the basic character that might be expected for an octahedral Fe–NH$_2$ species.$^{27}$ Therefore, this first $E_1$ redox event involves a square mechanism, with a preliminary $E_1C_1$ oxidation reaction (blue equations in Scheme 3.2) coupled to a second $E_2C_2$ reduction reaction (red equations in Scheme 3.2).

**Scheme 3.2.** Proposed square mechanism for the first redox event, $E_1$, at $0.4$ V. The presence of the TPA ligand is implied for the species shown.

We have further confirmed this mechanism by performing cyclic voltammetry at different scan rates, as depicted in Figure 3.5D. At low scan rates, where $C_1$ reaches equilibrium, two reductive peaks could be detected with similar intensities, corresponding to $[(TPA)Fe^{III}(NH_3)_2]^{3+}$ and $[(TPA)Fe^{III}(NH_3)(NH_2)]^{2+}$. However, when the scan rate was increased, the reduction peak at around $0.2$ V showed greater intensity as compared to the
peak at −0.4 V. This is a consequence of the relatively slow forward reaction, C, as compared to the fast scan rate, such that C does not reach equilibrium, and hence [(TPA)Fe(III)(NH$_3$)$_2$]$^{2+}$ is the main species reduced at the electrode.

Analysis of the peak currents at $E_1$ obtained from CVs at different scan rates reveals a linear relationship with $v^{1/2}$, following the predicted behavior from the Randles-Sevcik equation for an electrochemically reversible electron transfer process of a freely diffusing molecule in solution. Further analysis of the current corresponding to the oxidative peak with increasing concentrations of [(TPA)Fe(L)$_2$]$^{2+}$ again reveals the linear dependence expected for a one electron oxidation. This data, in combination with the well-behaved shift in $E_1$ potential with both NH$_3$ and NH$_4^+$ concentration, collectively support a primary process at $E_1$ to form a reactive Fe$^{III}$–NH$_2$ species.

3.2.4 Mechanistic insight into the catalytic process ($E_2$). We performed a similar analysis of the $E_2$ ($E_{cat}$) event where fast catalysis is observed. While a detailed analysis for this six-electron/six-proton process can in principle be performed, in practice such an analysis quickly becomes intractable. A common mechanistic approximation is therefore to consider the simplest case scenario, where electron transfer from the catalyst to the electrode is

Scheme 3.3. A simplified catalytic mechanism for ammonia oxidation at $E_2 \approx 1.1$ V. P and Q represent the inferred intermediates [(TPA)Fe$^{III}$(NH$_3$)(NH$_2$)$_2$]$^{2+}$ and its one-electron oxidized species [Fe$^{IV}$(NH$_3$)(NH$_2$)]$^{3+}$, respectively. The latter intermediate would then react with ammonia in a process that ultimately releases N$_2$. $E_{P/Q}$ is the potential for the P/Q redox couple, and $k_{obs}$ is the apparent second-order rate constant of the catalytic chemical step.
followed by a rate determining homogeneous catalytic reaction with the substrate, EC$_{\text{cat}}$, as shown in Scheme 3.3.$^{25,30}$

A related strategy has been previously applied to electrocatalytic redox processes for CO$_2$ reduction$^{31}$ and water oxidation,$^{32}$ and provides essential information regarding the mechanism and the overall kinetics for the catalytic process. Such an approach is justified here because the potential of the first EC step ($E_1$) is cathodically well separated from $E_2$. Furthermore, oxidation following an N–N bond formation step is expected to occur at less oxidizing potentials. In support of this notion, we found that a CV of [(TPA)Fe(L)$_2$]$^{2+}$ in acetonitrile with added hydrazine displayed a catalytic oxidation around 0.4 V. Our primary assumption is thus that electrocatalytic NH$_3$ oxidation to N$_2$ is triggered via a single electron transfer step that occurs at a substantially more oxidizing potential than all other steps. If correct, simplification to an EC mechanism is reasonable, and it then becomes possible to obtain a kinetic constant ($k_{\text{obs}}$) that reflects the overall rate of the catalytic reaction, after scaling for the number of electrons transferred (n).$^{33}$

To calculate $k_{\text{obs}}$, we performed a foot of the wave analysis (FOWA).$^{31}$ Due to non-Nernstian behavior in the ET step (see Supporting Information for further details), the Butler-Volmer law, which includes a charge transfer coefficient ($\alpha$) and the rate constant for interfacial electron transfer ($k_\text{s}$), was utilized.$^{34}$
Plotting the FIT equation versus $1/(1+\exp[F/(RT)(E_{\text{cat}}-E)])$ gives the value for pseudo first-order constant $k'_{\text{obs}} = k_{\text{obs}}[\text{NH}_3]$ (see SI). To use these equations, $\alpha$ and $k_s$ were determined from the precatalytic wave by analyzing the peak potential at different scan rates (see SI). We found an $\alpha$ value of 0.66, close to the typical value when use of the Butler-Volmer equation is necessary, and a diffusion coefficient $D_{\text{cat}}$: $9.5 \cdot 10^{-10}$ (m$^2$·s$^{-1}$), which is typical for molecular complexes in solution (see Supporting Information). With these values, we could obtain the kinetic constant for the electron transfer $k_s$ of around 111 m·s$^{-1}$. These data enable a FOWA according to Equation 3.3, which provides a second order apparent rate constant, $k_{\text{obs}}$, of $3.7 \cdot 10^7$ M$^{-1}$·s$^{-1}$ on average for different concentrations of catalyst (Figure 3.6A; see SI for details). Hence, this iron catalyst for AO is able to operate at a remarkably fast rate under a large applied bias of 1.1 V.

In addition to providing kinetic information, this method of analysis also offers important mechanistic information about the catalytic process, because the validity of these equations is intrinsically related to the validity of the assumed mechanism. Our assumed mechanism includes a rate-determining step which is first order both in catalyst and in NH$_3$. Accordingly, the calculated pseudo-first order rate constant, $k'_{\text{obs}}$, should be constant with respect to the [((TPA)Fe(L)$_2$)$_2$]$^{2+}$ concentration (because $k'_{\text{obs}}$ is normalized by its molarity), and should respond linearly to the NH$_3$ concentration. We have confirmed both relationships by analysis at different concentrations of [((TPA)Fe(L)$_2$)$_2$]$^{2+}$ and [NH$_3$] (Figure 3.6A,B).
3.3 Conclusion

With interest growing in the study of molecular (electro)catalysts for AO, a fascinating multi-electron redox reaction that represents the microscopic reverse of $\text{N}_2$-to-$\text{NH}_3$ conversion, electrochemically well-defined model systems are needed. In this context we have described in detail the capacity of a polypyridyl iron catalyst, $\left[(\text{TPA})\text{Fe(MeCN)}_2\right]^{2+}$, to perform AO at extremely fast rates ($\sim10^7\text{ M}^{-1}\cdot\text{s}^{-1}$ via FOWA) under the application of a 1.1 V applied bias. We have also used CPC to confirm that $\text{N}_2$ is selectively formed via this AO reaction, confirming as many as 16 eq of $\text{N}_2$ (32 eq $\text{NH}_3$ being consumed) per Fe. While TONs do not necessarily reflect overall efficiency of a catalyst, the value we measure is higher than other values reported to date. Mechanistic data extracted from a range of electrochemical studies suggest that an $\text{Fe}^{\text{III}}$–$\text{NH}_2$ species is generated at the first $E_1$ process (0.4 V vs Fc/Fc$^+$) via net H-atom removal from intermediate $\left[(\text{TPA})\text{Fe(NH}_3)_2\right]^{2+}$.
The first wave, $E_1$ at ~0.4 V vs Fc/Fc$, has been determined to correspond to a subsequent electron and proton transfer (EC) from a newly characterized $[(TPA)Fe(NH_3)_2]^{2+}$ species to generate a reactive terminal amide $[(TPA)Fe^{III}(NH_2)(NH_3)]^{2+}$ intermediate. This intermediate then undergoes a subsequent EC step at $E_2$ around 1.1 V, and this is the feature associated with catalytic AO. We speculate that at this potential Fe(III) is oxidized to Fe(IV), possibly via an $[(TPA)Fe^{IV}(NH_3)(NH_2)]^{3+}$ intermediate. Future studies will focus on the direct spectroscopic detection of these (or other) candidate intermediates. Regardless, we intuit that N–N bond forming steps occur subsequent to the generation of $[(TPA)Fe^{IV}(NH_3)(NH_2)]^{3+}$ or some similar species. We cannot yet determine from the available data whether or not such steps are intramolecular (e.g., via N–N coupling from adjacent Fe–NH$_x$ ligands) or intermolecular (e.g., via nucleophilic attack of NH$_3$ on Fe–NH$_x$). We are, however, able to say from FOWA that the overall catalytic reaction is first order in both $[(TPA)Fe(L)_2]^{2+}$ and NH$_3$.

The catalysis is ultimately arrested due to passivation of the BDD working electrode surface, likely via a combination of incorporation of nitrogen and iron, which can be detected via analysis of the electrode post CPC. It may be that exploring less labile polydentate ligands, such as those possessing a negative charge, will attenuate the lability of TPA that is operative in this system and ultimately leads to catalyst degradation. Such approaches may also afford a catalyst that operates at less oxidizing potentials, though this may in turn attenuate the overall catalytic rate.
3.4 References


9 Buhr, J. D.; Taube, H. Oxidation of $[\text{Os(NH}_3)_5\text{CO}]^{2+}$ to $[(\text{Os(NH}_3)_4\text{CO}_2\text{N}_2)]^{4+}$. *Inorg. Chem.*, 1979, 18, 2208–2212.


17 For select examples see: (a) Fillol, J. L.; Codolà, Z.; Garcia-Bosch, I.; Gómez, L.; Pla, J. J.; Costas, M. Efficient Water Oxidation Catalysts Based on Readily Available Iron Coordination Complexes. *Nat. Chem.*, 2011, 3, 807–813. (b) Zhang, B.; Li, F.; Yu, F.; Cui, H.; Zhou, X.; Li, H.; Wang, Y.; Sun, L. Homogeneous Oxidation of Water by Iron...


19 Charge passed and Faradaic efficiencies are averaged over six experiments. Turn over numbers are averaged over two experiments with 2.5 mmol [(TPA)Fe(MeCN)]OTf2, the optimum condition studied. Reported values for N2 are all corrected for background nitrogen, determined by integrating the oxygen peak in the gas chromatograms. For N2 produced by 2.5 mmol [(TPA)Fe(L)2]+, 1.0 equivalent of background N2 produced by the electrode has been subtracted.


23 The bis-ammine has a redox potential 300 mV lower than the bis-acetonitrile complex, therefore replacement of each MeCN by NH3 can be estimated to shift the potential by 150 mV assuming a linear relationship.


