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Facile conversion of ammonia to a nitride in a rhenium system that cleaves dinitrogen†

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Rhenium complexes with aliphatic PNP pincer ligands have been shown to be capable of reductive N₂ splitting to nitride complexes. However, the conversion of the resulting nitride to ammonia has not been observed. Here, the thermodynamics and mechanism of the hypothetical N–H bond forming steps are evaluated through the reverse reaction, conversion of ammonia to the nitride complex. Depending on the conditions, treatment of a rhenium(III) precursor with ammonia gives either a bis(amine) complex [(PNP)Re(NH₂)₂Cl]⁺, or results in dehydrohalogenation to the rhenium(III) amido complex, (PNP)Re(NH₂)Cl. The N–H hydrogen atoms in this amido complex can be abstracted by PCET reagents which implies that they are quite weak. Calorimetric measurements show that the average bond dissociation enthalpy of the two amido N–H bonds is 57 kcal mol^{−1}, while DFT computations indicate a substantially weaker N–H bond of the putative rhenium(IV)-imide intermediate (BDE = 38 kcal mol^{−1}). Our analysis demonstrates that addition of the first H atom to the nitride complex is a thermochemical bottleneck for NH₃ generation.

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Introduction

The interconversion of N₂ and NH₃ is important in fields that range from agriculture to sustainable energy.^{1,2} The heavy use of NH₃ in fertilizer manufacturing has resulted in an extensive global infrastructure for its transportation and storage.³ Coupled with its high energy density, this makes NH₃ an excellent candidate for a carbon-free chemical fuel, either through combustion or direct ammonia fuel cells (DAFCs).^{4–7} In order to realize this potential, it is beneficial to understand the individual steps of N–N and N–H bond formation and cleavage. One promising route to form the N–H bonds in NH₃ from N₂ is proton-coupled electron transfer (PCET).^{8a,9} Photo- or electrochemical energy may provide the necessary driving force for PCET-assisted N₂ reduction using water as a source of protons and electrons, thus providing a sustainable strategy for converting N₂ to NH₃.^{10–16} A growing number of homogeneous systems catalytically achieve this difficult transformation utilizing PCET.^{17–31}

It is also important to understand the reverse reaction, NH₃ oxidation to form N₂. One application of this reaction is for

releasing the chemical energy stored in N–H bonds for DAFC applications.^{6,7} In addition, the individual steps in NH₃ oxidation to N₂ are often the microscopic reverse of those used for PCET reduction of N₂ and thus help to elucidate potential mechanisms for PCET-assisted reduction of N₂ to NH₃.³² In this context, it is relevant that many examples of chemical N–H bond oxidation from NH₃-derived metal ammines yield metal nitride complexes.^{33–46} These systems utilize either chemical oxidants under basic conditions or H-atom abstracting (HAA) reagents for the ammine-to-nitride transformations.^{8b} In some systems, electrochemical oxidation of ammine complexes yields metal-nitride products.^{41,42,44} Other systems can generate N₂ as a product from the oxidation of NH₃-derived ammine complexes, either through chemical^{36,40,47–49} or electrochemical^{49–52} methods. These include recently reported homogeneous systems that catalytically form N₂ from NH₃ through both chemical^{43–45} or electrochemical^{44,53–55} N–H bond oxidation. N–N bond formation can occur *via* bimetallic N–N coupling (*e.g.*, between metal–NH_x species or metal nitrides)^{40,44,45} or nucleophilic attack on a metal–NH_x intermediate by NH₃.^{43,53,54,56}

Here, we study NH₃ oxidation in a well-defined system that is also capable of reductive functionalization of N₂ *via* an N₂-cleavage mechanism.^{57,58} Electrochemical reduction of (PNP)ReCl₂ (**1**, PNP = N((CH₂CH₂)P^tBu₂)₂) cleaves N₂ to form the nitride complex (PNP)Re(N)Cl (**2**), which contains a nucleophilic nitride ligand (Scheme 1, black arrow).^{59–61} This nitride can be alkylated and reduced to give N–C containing products,^{62,63} but PCET reduction of the nitride in **2** to form NH₃

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Scheme 1 Cycles that represent reductive N_2 splitting by (PNP)Re and PCET nitride reduction (gray cycle, not observed) or NH_3 oxidation (blue cycle, studied here).

(Scheme 1, grey arrows) was not observed because pincer protonation occurs rather than nitride protonation. Additional challenges are that the high energy of the lowest unoccupied molecular orbital (LUMO) of **2** prevents a reduction-first pathway, and that **2** is unreactive towards organic hydrogen-atom transfer (HAT) reagents or H_2 .⁵⁹ In this manuscript, we evaluate the reverse reactions (Scheme 1, blue arrows) to elucidate the factors that prevent PCET nitride reduction in this system. This fundamental information may help to improve NH_3 oxidation catalysis and to avoid bottlenecks in NH_3 generation by future N_2 -cleaving systems, and importantly provides a thermochemical framework for nitrogen fixation products beyond ammonia.

Results

Binding and deprotonation of NH_3

Introduction of 1 atm of NH_3 gas to a solution of the dichloride complex **1** in benzene- d_6 or tetrahydrofuran- d_8 (THF- d_8) results in an immediate color change from purple to brown. 1H NMR spectroscopy reveals the formation of a new C_s -symmetric product **3** in >95% yield (Scheme 2, top). The chemical shifts of **3** (Fig. S1†) are characteristic of a non-magnetic ground state non-, and the lack of noticeable temperature-independent paramagnetism, which is often observed in Re^{III} complexes,^{64,65} suggests that the two strongly π -donating amide ligands sufficiently destabilize the spin triplet state to give a well-isolated singlet ground state. A notable 1H resonance integrating to 2H is found at δ 12.7 ppm. A 1H - ^{15}N HSQC spectrum of a natural-abundance sample shows a ^{15}N cross-peak from this resonance at δ -260 ppm (Fig. S2†), confirming that it corresponds to protons bound to N. This ^{15}N chemical



Scheme 2 Reactivity of **1** with N_2 and NH_3 .

shift is significantly upfield from related nitride complexes (371–393 ppm) and closer to that for the protonated PNP backbone of $[(^1H)PNP]Re(N)Cl]^+$ (**5**) (–336 ppm).⁶⁶ All spectroscopic signatures are consistent with the formulation of **3** as the amido complex (PNP)Re(NH_2)Cl.

On a preparative scale, addition of 1 atm NH_3 to a solution of **1** gives **3** as the major product, which is isolated from the reaction in 61% yield. The solid-state structure of **3** was elucidated *via* single crystal X-ray diffraction (XRD) and the N-bound hydrogen atoms were located in the Fourier map (Fig. 1). The Re– NH_2 bond in **3** is 0.3 Å longer than the Re≡N bond in the nitride complex **2** (Table 1).⁶⁶ In the supporting ligand, the (PNP)–Re bond is 0.1 Å shorter in **3** than in the Re–nitride complex **2**, indicating increased π -bonding from the nitrogen of the pincer ligand in **3** (Table 1). The (PNP)–Re and Re– NH_2 amide bond lengths in **3** are within 0.02 Å of each other with

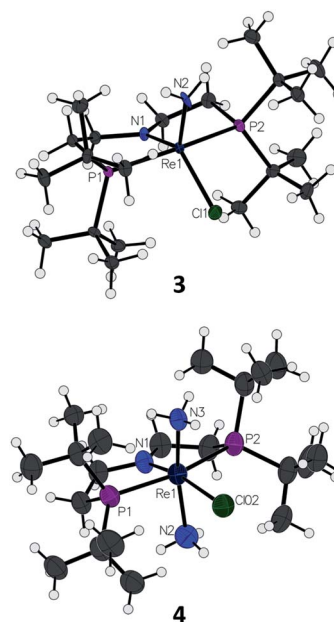


Fig. 1 Solid-state structures of Re–amide complex **3** and $Re(NH_3)_2$ complex **4** ($BARF_4$ ion omitted) with thermal ellipsoids at 50% probability.



Table 1 Selected bond lengths (Å) and bond angles (°) of complexes 2–4

Bond/angle	2	3	4
Re1–N1	2.033(6)	1.936(3)	1.894(5)
Re1–N2	1.643(6)	1.959(3)	2.150(5)
Re1–N3	—	—	2.193(6)
Re1–Cl1	2.441(2)	2.384(1)	2.495(2)
Re1–P1	2.443(2)	2.397(1)	2.424(2)
Re1–P2	2.435(2)	2.382(1)	2.425(2)
N1–Re1–N2	105.8(3)	115.5(1)	84.8(2)
N1–Re1–N3	—	—	165.6(2)
N2–Re1–N3	—	—	109.6(2)
N1–Re1–Cl1	106.5(2)	108.8(1)	83.4(1)
N2–Re1–Cl1	147.7(2)	135.6(1)	167.0(2)
N1–Re1–P1	100.4(2)	95.1(1)	91.0(1)
N1–Re1–P2	99.9(2)	95.5(1)	90.5(1)

planar coordination of the nitrogen atoms in both cases ($\Sigma_{\text{PNP}} = 360^\circ$, $\Sigma_{\text{NH}_2} = 357^\circ$). The PNP and NH_2 amides are oriented to π -donate into the same Re d orbital, which gives modest lengthening (0.04 Å) of the (PNP)–Re bond in **3** compared to the Re-dichloride complex **1**.⁵⁹ This also likely contributes to increased pyramidalization of the dialkylamide group ($\Sigma_{\text{PNP}} = 348^\circ$) in **2**. These structural differences are accompanied by a change of the rhenium coordination geometry from square pyramidal ($\tau_5 = 0.14$) in complex **2**, in which the coordination site *trans* to the nitride is open, toward trigonal bipyramidal ($\tau_5 = 0.48$) in complex **3**.

When 1 equiv. of NH_3 gas was added to a solution of **1** in THF- d_8 at -80°C , ^1H and $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of the reaction showed a mixture of diamagnetic products (Fig. 2, middle). Addition of another 4 equiv. of NH_3 gas (for a total of 5 equiv. NH_3 per Re) resulted in full consumption of **1** and observation of **3** in 71% yield (Fig. 2, top). It is likely that dehydrohalogenation of the putative intermediate $(\text{PNP})\text{Re}(\text{NH}_3)_2\text{Cl}_2$ by NH_3 to form NH_4Cl is required to drive the formation of **3**. This implies that coordination to Re^{III} significantly increases the acidity of the N-bound protons.⁶⁷

Addition of 1 atm NH_3 to a solution of **1** containing an equivalent of $\text{NaBar}^{\text{F}}_4$ ($\text{Ar}^{\text{F}} = 3,5\text{-bis(trifluoromethyl)phenyl}$) in

THF- d_8 at -80°C resulted in a color change from purple to light green and formation of the new diamagnetic complex **4** by ^1H and $^{31}\text{P}\{^1\text{H}\}$ NMR spectroscopy (see Scheme 2). In contrast to **3**, complex **4** exhibits C_{2v} symmetry, broadened resonances, and a new peak at $\delta = 5.47$ ppm that integrates to 6H (Fig. S3†). Despite no identifiable cross-peaks in the ^1H – ^{15}N HSQC spectrum of **4**, N–H stretching bands were observed in the infrared (IR) spectrum at 3392, 3353, 3245, and 3174 cm^{-1} . The molecular structure of **4** in the solid state shows the six-coordinate, cationic bis-ammine adduct $[(\text{PNP})\text{Re}(\text{NH}_3)_2\text{Cl}][\text{Bar}^{\text{F}}_4]$ with a distorted octahedral geometry (Fig. 1). In comparison to **3**, complex **4** shows lengthened Re–N bonds (2.172(5) Å vs. 1.959(3) Å) due to the lack of π -donation. With no strong π -donor ligands to compete with π -donation from the PNP amide, **4** contains a Re–PNP bond distance that is shorter than in **3** and **2** (Table 1). The flexibility of the PNP–Re interaction to accommodate the changes in ligand donor characteristics from ammine to nitride is also evident from the change in the PNP–Re bond lengths and PNP pyramidalization from **2**–**4**.

Reactivity of $[(\text{PNP})\text{Re}(\text{NH}_3)_2\text{Cl}]^+$

To assess the plausibility of an ammine complex as an intermediate during formation of **3**, a solution of **4** in THF- d_8 was treated with 1 atm of NH_3 , which gave no reaction. However, addition of a slight excess of potassium hexamethyldisilazide (KHMDs) (Scheme 3) caused an immediate color change and formation of **3** as the major product in 61% yield, as judged by ^1H and $^{31}\text{P}\{^1\text{H}\}$ spectroscopy (Fig. S4†).

Cyclic voltammetry (CV) of **4** in THF under Ar shows irreversible redox processes, a reduction at $E_{\text{pc}} = -1.95$ V vs. $\text{Cp}_2\text{Fe}^{+/0}$ and an oxidation at $E_{\text{pa}} = -0.58$ V vs. $\text{Cp}_2\text{Fe}^{+/0}$ (Fig. S16†). The position of the reduction peak is similar to the reversible reduction of **1** under Ar at -2.00 V vs. $\text{Cp}_2\text{Fe}^{+/0}$.⁶⁸ The first reduction of **1** under Ar was previously attributed to the formation of $[(\text{PNP})\text{Re}^{\text{II}}\text{Cl}_2]^-$, which is followed by chloride dissociation to form $(\text{PNP})\text{Re}^{\text{II}}\text{Cl}$ which is subsequently reduced again.⁶⁰ The difficulty of reducing **4** suggests that it is quite electron-rich despite its positive charge, but the lack of reversibility prevents further interpretation.

In an attempt to assess the species formed upon reduction, **4** was treated with a chemical reductant. Addition of 1.2 equiv. CoCp^*_2 to a solution of **4** in THF- d_8 under N_2 gave complete consumption of **4** but the Re amide **3** was formed (Fig. S5†). The spectroscopic yield of **3** was only 60%. The fate of the lost proton and electron in the formation of **3** remain unknown. Analysis of



Fig. 2 ^1H and $^{31}\text{P}\{^1\text{H}\}$ NMR spectra of **1** in THF- d_8 without NH_3 (bottom, maroon), with 1 equiv. NH_3 added (middle, green), and with 5 equiv. NH_3 added (top, blue). Spectroscopic yields reported vs. PMe_3 in a capillary.



Scheme 3 Reactivity of **4** with stoichiometric base or reductant.

the headspace following the reaction showed no detectable amount of H_2 (<1% yield).

N–H abstraction from Re–amide complex (PNP)Re(NH₂)Cl

We hypothesized that abstraction of H atoms from **3** would lead to the nitride (Scheme 1, blue), by analogy with other reported systems.^{37–40,43,45,46} In the following, we assume that formal H[•] abstraction by the hydrogen atom abstraction (HAA) reagents is most likely concerted, based on the known difficulty of stepwise PCET pathways.⁸ Addition of 2 equiv. of either 2,4,6-*tert*-butylphenoxyl radical (^tBu₃PhO[•]) or 2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPO[•]) as HAA reagents to a solution of **3** in THF-*d*₈ or benzene-*d*₆ at ambient temperature gives rapid and quantitative (>99%) formation of **2** (Scheme 4), as judged by ¹H and ³¹P{¹H} NMR spectroscopy (Fig. S6†). This is accompanied by the formation of 2 equiv. of ^tBu₃PhOH or TEMPOH. These reagents have O–H bond dissociation free energies (BDFE_{O–H}) of 74.4 and 65.5 kcal mol^{–1} in THF, respectively.^{8b,69} When **3** is mixed with only 1 equiv. of TEMPO[•], only half of **3** is consumed, showing that the second H-atom abstraction is more favorable than the first (Fig. S7†). The absence of reactivity of **2** with excess ^tBu₃PhOH or TEMPOH supports this notion.

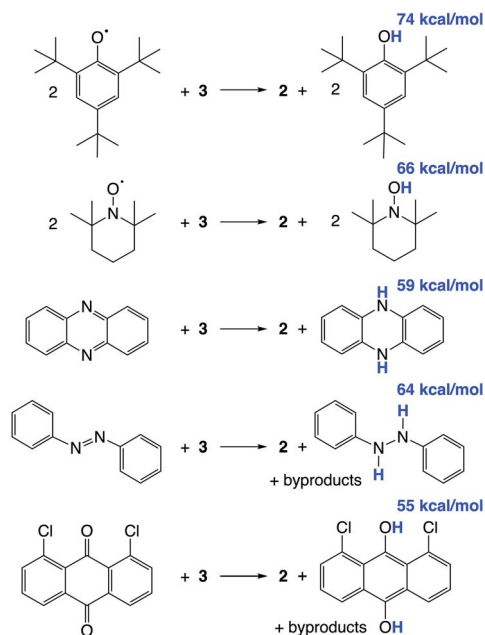
Additional HAA reagents were used to further bracket the N–H bond strengths (Scheme 4). While **3** did not react with 5,10-phenazine (5 equiv) in THF-*d*₈ at ambient temperature, heating to 80 °C gave quantitative (>98%) conversion to **2** and 5,10-dihydrophenazine (average BDFE_{N–H} = 58.7 kcal mol^{–1} in MeCN⁶⁹) after 21 h (Fig. S8†). Accordingly, no reaction was observed between **2** and 10 equiv. of 5,10-dihydrophenazine even after prolonged heating at 80 °C. Oxidation of **3** to form **2** was also observed when using 1.5 equiv. of azobenzene (54% yield of **2** after 72 h at 60 °C) and 1,8-dichloro-9,10-

anthraquinone (67% yield of **2** after 4 d at ambient temperature). ¹H NMR spectra of reaction mixtures showed the formation of 1,2-diphenylhydrazine and 1,8-dichloro-9,10-anthracenediol (average BDFE_{X–H} = 60.9 [in MeCN] and 55.4 kcal mol^{–1} [THF]), respectively.⁶⁹ However, these reactions form multiple products, so quantitative thermochemical information cannot be derived from the product formation in these cases. Quantification of the PCET thermochemistry was therefore carried out by titration calorimetry as detailed below.

Stepwise ET–PT from (PNP)Re(NH₂)Cl

The reactivity of **3** with HAA reagents suggests that the N–H bonds in the amide ligand can be easily oxidized *via* concerted removal of an H-atom.⁸ We were also interested to determine whether the conversion to **2** is possible through stepwise PCET, with deprotonation and 1e[–] oxidation of **3**.^{41,42,44} In order to test a PT–ET (proton transfer followed by electron transfer) pathway, a solution of **3** was mixed with up to 12 equiv. of 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU, pK_a of conjugate acid = 16.9), phosphazene base P₁-tBu-tris(tetramethylene) (pK_a of conjugate acid = 20.2), or phosphazene base P₄-tBu (pK_a of conjugate acid = 33.9) in THF-*d*₈ at ambient temperature.⁷⁰ No reaction of **3** with any of these strong bases was observed by ¹H and ³¹P{¹H} NMR spectroscopy, indicating that the amide ligand in **3** is a poor Brønsted acid.

In other tests, we explored whether a stepwise ET–PT pathway (electron transfer followed by proton transfer) is feasible. CV of **3** in THF shows an irreversible oxidation wave ($E_{pa} = -0.61$ V vs. Cp₂Fe^{+/0}) at a scan rate of 100 mV s^{–1} (Fig. 3). However, increasing the scan rate to 1 V s^{–1} results in a distinct anodic shift of the oxidation event and increased reversibility, indicating chemical follow-up steps at a time-scale of the CV experiment. This potential is similar to those for the oxidation of both the dichloride complex **1** and the Re(NH₃)₂ complex **4**.⁶⁰ Further analysis of the CV has not been fruitful because of the lack of reversibility and formation of unknown byproducts (see below). However, electrolysis of a solution of **3** in the presence of 2,6-lutidine (pK_a = 7.2 in THF⁷⁰) at a potential of +0.6 V relative to the open circuit potential (OCP) resulted in steady passing of charge up to 2.2 equiv. e[–] (Fig. S15†) and a change in color from



Scheme 4 Reactivity of **3** with organic HAA reagents, with BDFE_{O–H} or average BDFE_{X–H} of the organic products in THF given in blue.



Fig. 3 Cyclic voltammogram of the first oxidation of **3** (0.2 mM) in 0.2 M NBu₄PF₆ solution in THF under N₂ using a glassy carbon working electrode, Pt wire auxiliary electrode, and Ag wire pseudoreference. Potentials referenced to Cp₂Fe^{+/0} after the experiments.

brown to orange. Rhenium(v) nitride complex **5**, in which the backbone is protonated,⁶⁶ was isolated from the post-electrolysis mixture in 69% isolated yield (Fig. S17†). The combination of removing an electron at -0.61 V and a proton with lutidine is thermodynamically equivalent to an “effective BDFE” of 56 kcal mol^{-1} ,^{69,71} so this e^-/H^+ removal is thermodynamically similar to the HAA reactions above.

In an effort to identify oxidation products of **3**, chemical oxidation was carried out with 1.1 equiv. of $[\text{Cp}_2\text{Fe}][\text{PF}_6]$ in $\text{THF}-d_8$ (Scheme 5). The major product identified from the resulting ^1H and $^{31}\text{P}\{^1\text{H}\}$ NMR spectra was **5** in 50% yield (Fig. S9†).⁶⁶ Furthermore, the Re^{III} -dichloride complex **1** was obtained in 25% yield, as well as a brown precipitate that could not be identified. The formation of both Re^{III} and Re^{V} complexes from the $1e^-$ oxidation of **3** implies disproportionation; however, these products are not formed in a 1 : 1 ratio, implicating additional decomposition pathways. The product mixture that can be identified spectroscopically does not account for all of the Re, N, or H atoms present in the starting material. To test whether the missing H atoms could be released as H_2 from weakened N–H bonds during the reaction, the reaction was repeated on a larger scale. Analysis of the THF -soluble products from the reaction showed formation of **1** and **5** in 17% and 52% yield, respectively, and no H_2 was detected from analysis of the headspace (<1% yield).

We also tested whether oxidation of **3** to a nitride could be facilitated by providing an exogenous base for the deprotonation steps and by using 2 equiv. of oxidant (Scheme 5). Accordingly, 20 equiv. 2,6-lutidine and 2.2 equiv. $[\text{Cp}_2\text{Fe}][\text{PF}_6]$ were added to a solution of **3** in $\text{THF}-d_8$, forming **5** in 82% spectroscopic yield, and no observable **1** (Fig. S10†). An unidentified brown solid was formed as a byproduct in both reactions, suggesting decomposition. The formation of unknown byproducts deterred us from further mechanistic analysis.

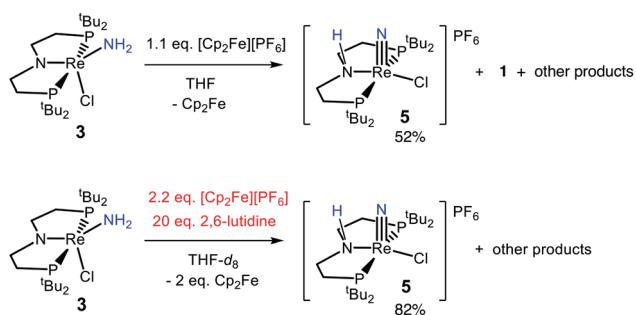
Calorimetric measurement and DFT calculations of N–H bond oxidation from $(\text{PNP})\text{Re}(\text{NH}_2)\text{Cl}$

Since the reaction of **3** with 2 equiv. $^t\text{Bu}_3\text{PhO}^\bullet$ to form **2** and 2 equiv. $^t\text{Bu}_3\text{PhOH}$ proceeds quantitatively, we chose this reaction for calorimetric determination of the reaction enthalpy.^{72,73} The

titration of **3** with $^t\text{Bu}_3\text{PhO}^\bullet$ in THF using isothermal titration calorimetry (ITC) results in an exotherm of $-50.9 \text{ kcal mol}^{-1}$ until 2.0 equiv. of $^t\text{Bu}_3\text{PhO}^\bullet$ are added (Fig. S18†).^{33,73} Thus, on average, each abstraction of an H atom from **3** gives an enthalpy change of $-25.4 \text{ kcal mol}^{-1}$. Since the bond dissociation enthalpy of $^t\text{Bu}_3\text{PhO}-\text{H}$ in THF is $80.8 \pm 1 \text{ kcal mol}^{-1}$,[‡] the average of the two $\text{BDE}_{\text{N-H}}$ values of **3** in THF is $55.4 \pm 1 \text{ kcal mol}^{-1}$.

DFT calculations were used to obtain more insight into the thermodynamics of each PCET step from amide complex **3** to nitride complex **2**. The B3LYP functional and def2-TZVP basis, together with standard solvent and dispersion corrections gave excellent agreement with the metrical parameters of the crystal structure of **3**, and predicted the redox potential for $3^{+/0}$ ($E = -0.65 \text{ V}$) close to the observed wave at -0.61 V in the experimental CV (see ESI†). Computation of the putative PCET intermediate confirmed that the $S = 1/2 \text{ Re}^{\text{IV}}$ -imide $(\text{PNP})\text{Re}(\text{NH})\text{Cl}$ (**LRe=NH**) is the most stable tautomer; an isomeric amido-rhenium(IV) complex, $(\text{PNP}^*)\text{Re}(\text{NH}_2)\text{Cl}$ ($\text{PNP}^* = \text{N}(\text{CHCH}_2\text{-P}^t\text{Bu}_2)(\text{CH}_2\text{CH}_2\text{P}^t\text{Bu}_2)$), with an unsaturated PNP backbone proved higher in free energy by 12 kcal mol^{-1} . The optimized structure of **LRe=NH** shows a strongly bent parent imido ligand with a computed $\text{Re}-\text{N}-\text{H}$ angle of 133° . Bending reduces the antibonding π -interaction of the multiply bonding imido ligand with the metal centred SOMO after reduction. In consequence, the Re–imide bond is considerably elongated (1.80 \AA) with respect to the parent nitride (DFT: 1.66 \AA).

Computations of the *enthalpies* associated with each sequential H-atom transfer from **3** to $^t\text{Bu}_3\text{PhO}^\bullet$ to give $^t\text{Bu}_3\text{PhOH}$ in THF gave an average calculated $\text{BDE}_{\text{N-H}}$ of 57 kcal mol^{-1} (Scheme 6).^{38,40,58} This is close to the calorimetrically determined average BDE of $55.4 \pm 1 \text{ kcal mol}^{-1}$. The *free energy* of conversion of **3** to **2** via H-atom transfer to $^t\text{Bu}_3\text{PhO}^\bullet$ gave an average calculated $\text{BDFE}_{\text{N-H}}$ in **3** of 51 kcal mol^{-1} . Further insight comes from the hypothetical $1e^-/1H^+$ steps. The $\text{BDFE}_{\text{N-H}}$ for removing the first H from the amide ligand in **3** was calculated to be 69 kcal mol^{-1} , which is substantially higher than the computed $\text{BDFE}_{\text{N-H}}$ of the second N–H bond (in **LRe=NH**), 33 kcal mol^{-1} . These computations show that the N–H bond in **LRe=NH** is particularly weak, which is consistent with both the facile, irreversible oxidation of **3** with HAA reagents and the inability to observe this putative parent imido complex (see ESI†).



Scheme 5 Reactions of **3** with chemical oxidants.



Scheme 6 DFT (B3LYP/def2-TZVP) computed thermochemistry for the oxidation of **3** to **2** via stepwise H-atom removal.

Discussion

NH₃ conversion to nitride with (PNP)Re

The conversion of NH₃ to a nitride is quite facile in this system. Using excess NH₃, complex **1** goes directly to the rhenium amide complex **3**. The intermediate Re^{III}-ammine complex, potentially (PNP)Re(NH₃)Cl₂, has been neither spectroscopically observed nor isolated, which we attribute to rapid dehydrohalogenation by free NH₃. This can be avoided through preparation of the cationic Re^{III}(NH₃)₂ complex **4**, which can be deprotonated to form isolable complex **3**. Further oxidation of **3** to Re–nitride complexes is also facile, forming **5** using either hydrogen atom abstracting (HAA) reagents or forming **5** *via* 2e[−] chemical oxidation in the presence of a weak base. Thus, the complete conversion of NH₃ to a nitride in this system is achievable in good yield using 2e[−] electrochemical oxidation in the presence of base (Scheme 7). From a functional standpoint, such facile formation of a nitride complex from NH₃ is attractive considering the mild oxidation potential used (−0.6 V) and the possibility for excess NH₃ to serve as an exogenous base.⁸¹ However, turnover to achieve catalytic NH₃ oxidation to N₂ would require a nitride coupling step.^{35,82,83} Although nitride coupling reactions between other late transition-metal nitrides bearing similar supporting pincer ligands have been reported,^{46,84–86} nitride coupling is not observed in this system due to the strong Re–nitride bond in complex **2**.⁶⁶ We attribute this to a thermodynamic difficulty because the reverse reaction, reductive N₂ cleavage to form **2**, is very exergonic.

The calorimetric titrations give an average bond enthalpy for the two N–H bonds in **3** of 55.4 ± 1 kcal mol^{−1} in THF. This is a rare example of an experimentally-derived bond energy in the context of NH₃ oxidation.⁷⁴ In contrast, almost all other literature values (Table 2; see also ref. 45 and 75) are estimated through bracketing experiments or computational models.³² The computationally derived average BDE_{N–H} of 57 kcal mol^{−1} agrees with the experimental value, and the computations indicate that the analogous average BDFE (free energy) is 51 kcal mol^{−1}. This value represents a substantial weakening from the BDFE_{N–H} of free NH₃ (100 kcal mol^{−1}).⁸

The oxidation of ammine complexes to form nitrides using HAA reagents is preceded in other systems using ^tBu₃PhO[•] as the oxidant (forming ^tBu₃PhOH).^{37–40,43,46} H-atom abstraction from the amide in **3** can be achieved with HAA reagents to form X–H bonds that are up to 15 kcal mol^{−1} weaker than the O–H bond in ^tBu₃PhOH, though the phenazine reaction requires heating. The first HAA from **3** using organic reagents can be thermodynamically unfavorable by up to 10 kcal mol^{−1} because

nitride formation is driven by the much more favorable second N–H oxidation.^{37,38,45}

Though we have no experimental evidence for the intermediate imido complex **LRe=NH**, we considered its properties obtained from a DFT model. These computations indicate that the N–H bond in **LRe=NH** is especially weak, with a BDFE of 33 kcal mol^{−1}. This bond is 36 kcal mol^{−1} weaker than the calculated first BDFE_{N–H} of amido complex **3**. Additionally, the N–H bond in **LRe=NH** is calculated to be 10 kcal mol^{−1} weaker than the 43 kcal mol^{−1} value computed for the closely related (PONOP)Re^{IV}=NH (PONOP = 2,6-bis-(diisopropylphosphinito)pyridine).⁵⁸

As this last example shows, the thermochemistry of N–H bond forming and breaking is of particular interest for understanding how to achieve efficient N₂ to NH₃ interconversion. Table 2 compares our experimental values to literature values, typically derived from computational modelling. Complex **3** and its analogue (PONOP)Re(NH₂)(Cl)₂, both pincer-supported Re^{III}–NH₂ complexes, have low BDFE_{N–H} values. Interestingly, the Re^{IV}=NH compounds exhibit weaker imide N–H bonds than those of other metals. Some of the weakest N–H bonds were calculated for Mo–NH intermediates in the Chatt⁷⁸ (37 kcal mol^{−1}) and Schrock⁷⁶ (42 kcal mol^{−1}) systems, which can undergo complete N₂ reduction to ammonia.⁷⁹ Consistent with earlier systems (Table 2), amide intermediates consistently exhibit stronger N–H bonds than their corresponding imide intermediates. However, the difference between these two bond energies is particularly large in the Re systems, especially **3** where ΔBDFE_{N–H} = 36 kcal mol^{−1}. These values can be qualitatively rationalized by the very strong Re- and Mo-nitride bonds that arise when a d² configuration is reached, and the less favorable M–N π bonding at higher d-electron counts.

Relevance to the PCET nitride reduction step of NRR

Nitride complex **2** is readily formed *via* electrochemical N₂ cleavage,⁶⁰ so the conversion of NH₃ to the nitride ligand in **2** represents part of the reverse pathway from N₂ to NH₃ (Scheme 1). This would involve N₂ cleavage to form **2** followed by 3e[−]/3H⁺ PCET reduction of the nitride. In a recent review, Chirik and coworkers highlighted the lack of data in the literature on the bond strengths of N–H bonds in NH₂ and NH complexes in systems that perform N₂ reduction,⁷⁶ and the studies here are an important step toward understanding these species quantitatively. The thermochemical data from this study identify specific challenges associated with steps during the conversion of N₂ to NH₃.

One clear challenge in the PCET reduction of **2** is formation of the first N–H bond, which would give a very weak bond in **LRe=NH** with a BDFE_{N–H} of only 33 kcal mol^{−1}.^{76,78,79} One approach that has been used to overcome this difficulty in literature systems is the use of potent acid/reductant pairs to form the weak N–H bonds,^{21,29,30,58,79,87} though this is complicated in the current system by the ease of protonation of the pincer amide group.⁶⁶ In general, the instability of the imide species is identified as a key hindrance, because the imide intermediate must be accessed, even transiently, on the way to



Scheme 7 Full conversion of NH₃ by **1** to a nitride in complex **5**.



Table 2 Comparison of the measured average N–H bond dissociation enthalpy in **3** to computed N–H bond dissociation free energies (BDFE) of NH_x ligands in other relevant systems

Complex	Solvent	BDFE _{N–H} (kcal mol ^{−1})			Reference
		NH ₃	NH ₂	NH	
(PNP)Re(NH _x)Cl (3 , computed)	THF	—	69	33	This work
(PNP)Re(NH _x)Cl (3 , experimental)	THF	—	57(ave) ^a	—	This work
<i>cis</i> -(PONOP)Re(NH _x)Cl ₂	THF	—	78	43	58
(PNP)Ir(NH _x)	Gas phase	—	95 ^a	71 ^a	46
<i>trans</i> -[(Ph-tpy)(PPh ₂ Me) ₂ Mo(NH _x)] ⁺	THF	46	64	—	74 and 75
<i>cis</i> -[(Cp)(P ^{Ph} ₂ N ^{tBu} ₂)Mo(NH _x)(CO)] ⁺	Et ₂ O	84	61	—	38
[(PY5)Mo(NH _x)] ²⁺	MeCN	68	65	64	40
[(Cp*)(P ^{tBu} ₂ N ^{Ph} ₂)Ru(NH _x)] ⁺	THF	83	89	72	43
[(tpy)(^{NMe2} bpy)Ru(NH _x)] ²⁺	THF	79	86	—	56
(TMP)Ru(NH _x) ₂	C ₆ H ₆	82	93	75	45
[(tpy)(^{NMe2} bpy)Fe(NH _x)] ²⁺	THF	82	90	—	56
[(^{Ph} NCH ₂ CH ₂) ₃ N]Mo(NH _x)	—	52	64	42	76
[(BP ₃)Fe(NH _x)] ⁺	Et ₂ O	—	80	65	77
(F)(H ₂ PCH ₂ CH ₂ PH ₂) ₂ Mo(NH _x)	Benzene	41	92	37	78
(salen)Mn(NH _x)	Gas phase	85	84	60	79
(η ⁵ -C ₅ Me ₄ SiMe ₃) ₂ Ti(NH _x)	Gas phase	42	79	—	80

^a Bond dissociation enthalpies (BDE).

subsequent reduction that gives the amide species. The low driving force for PCET-assisted reduction of nitride **2** can be attributed to an overstabilization that results from strong Re≡N triple bonding.

Besides these thermochemical considerations, an encouraging result is that the amide complex **3** can be readily oxidized even though the first step (formation of **Re=NH**) is uphill. The conversion of **3** to **2** by TEMPO, for instance, occurs within minutes even though the first step is thermodynamically uphill by 3 kcal mol^{−1}. The ability of the first H⁺ abstraction to proceed indicates that the barriers for the H-atom transfer reactions are low. Therefore, even less stable imide intermediates may be sufficient to allow rapid catalysis in the future.

Conclusions

Rhenium-amide and -ammine complexes have been isolated, and they are readily oxidized to the corresponding metal-nitride complex. The conversion of NH₃ to a nitride by Re(PNP)Cl₂ (**1**) to form Re(PNP)(N)Cl (**2**) represents the first example of an NH₃-to-nitride transformation at Re. Since **2** can also be generated *via* electrochemical N₂ cleavage, this system is relevant both to N₂ reduction and ammonia oxidation. Facile, initial deprotonation of NH₃ occurs upon coordination to **1** to form the amide complex Re(PNP)(NH₂)Cl (**3**). The subsequent conversion to Re≡N can be accomplished with weak hydrogen atom acceptors that provide low driving force for the H⁺ transfer, indicating that the reactions are kinetically facile.

Calorimetric measurements of the conversion of **3** to **2** with ^tBu₃PhO⁺ show that the average BDE_{N–H} of **3** in THF is 55.4 ± 1 kcal mol^{−1}. This is a rare experimental thermochemical measurement of N–H bond strengths relevant to N₂/NH₃ interconversions.⁷⁶ Computations show that this average is derived from two disparate N–H bond strengths, as the second

N–H BDE in **3** (77 kcal mol^{−1}) is much stronger than the N–H BDE in the putative imide intermediate **LRe^{IV}=NH** (38 kcal mol^{−1}). The ability to form the weak N–H bond in this imide, even transiently, is identified as a crucial bottleneck for N₂ to NH₃ conversion in this system. The combination of bracketing,⁴⁶ calorimetry and DFT is a powerful strategy that will continue to elucidate the steps of N₂ reduction at metal complexes.

Data availability

Crystallographic data have been deposited at the CCDC with deposition numbers 2100974-2100975 and can be obtained from <https://www.ccdc.cam.ac.uk/structures/>. Other data supporting this article have been uploaded as part of the ESI.†

Author contributions

Conceptualization: G. P. C., J. M. M., P. L. H.; investigation and analysis: G. P. C., D. D., J. E. W., B. Q. M., J. B. C.; writing, interpretation: G. P. C., D. D., J. E. W., S. S., J. M. M., P. L. H.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

‡ The BDFE of $t\text{Bu}_3\text{PhO-H}$ in THF has recently been reported as $74.4 \text{ kcal mol}^{-1}$,⁶⁹ so the bond dissociation enthalpy (BDE) can be estimated as $80.7 \text{ kcal mol}^{-1}$. This takes TS° for H^\bullet in THF to be $6.3 \pm 0.2 \text{ kcal mol}^{-1}$, the mean of the $TS^\circ(\text{H}^\bullet)$ values for moderately polar aprotic solvents.

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