Metal-Ligand Cooperative Synthesis of Benzonitrile via Electrochemical Reduction and Photolytic Splitting of Dinitrogen


Dedicated to Prof. Dr. F. Tuczek on the occasion of his 60th birthday

Abstract: Thermal nitrogen fixation relies on strong reductants to overcome the extraordinarily large N–N bond energy. Photochemical strategies that drive N₂ fixation are scarcely developed. Here, the synthesis of a dinuclear N₂-bridged complex is presented upon reduction of a rhenium(III) pincer platform. Photochemical splitting into terminal nitride complexes is triggered by visible light. Clean N₂-transfer with benzoyl chloride to free benzamide and benzonitrile is enabled by cooperative 2H/2e⁻ transfer of the pincer ligand. A three-step cycle is demonstrated for N₂ to nitrile fixation that relies on electrochemical reduction, photochemical N₂-splitting and thermal N₂-transfer.

The Haber-Bosch process consumes large amounts of energy, for the generation of H₂. Electrocatalytic N₂ reduction has therefore been targeted as an attractive alternative. Nitrogen fixation at ambient conditions with molecular catalysts has seen remarkable progress, in some cases also for electrochemical ammonia generation. The direct synthesis of other compounds than NH₃ from N₂ remains a formidable challenge. Catalytic protocols are only known for trisilylamine, Nitrides, isocyanates, silylamines, and borylamines have been synthesized within stoichiometric, cyclic reaction sequences that allow for evaluating strategies to offset the extremely strong N–N bond (225 kcal/mol), enable E–N (E = C, Si, B) bond formation and deliver six reduction equivalents. All reported ‘synthetic cycles’ proceed through initial N₂ cleavage into nitride complexes. Subsequent N₂-transfer typically requires strong electrophiles like alkyl triflates. The thermochemistry of N₂ splitting must therefore be tuned to avoid nitride overstabilization and enable functionalization with reagents that are more compatible with reductive conditions.

We have examined N₂ activation and splitting, i.e. triggered by chemical or electrochemical reduction of pincer halide complexes. The rhenium(III) precursor [ReCl₂(PNP⁵Bu₁₂)] (1; PNP⁵Bu₁₂ = N(CH₂CH₂PBu₂)₂) exhibits a complex mechanism via rapid Re⁺/Re⁰-reduction, N₂-binding, halide loss, Re⁺/Re⁰-reduction and Re⁰/Re⁵-comproportionation. Splitting of the resulting dinuclear complex [(µ-N₂)(ReCl(PNP⁵Bu₁₂)] (2) is rate-determining and gives the nitride complex [ReNCI(PNP⁵Bu₁₂)] (3). A simple electronic structure model for the Re–N–N–Re core of key intermediate 2 (π¹c₁₂² configuration; Scheme 1) provides a starting point to tune the thermochemistry. We now report a modified platform that splits N₂ photolytically into more reactive nitrides that can be transferred with benzoyl chloride. The pincer ligand serves as 2e⁻/2H⁺ reservoir, enabling electrochemical N₂ reduction, photochemical splitting and thermal transfer within a three-step cycle.

The Re³ complex [ReCl₃(HNP⁵P₅)] (4) was obtained in 70% isolated yield. In contrast to 1, the amide complex [ReCl₃(PNP⁵P₅)] could not be isolated in analytical purity. 4 displays strongly shifted NMR resonances, as exemplified by the ⁳¹P(H)-NMR signal (δ = -1525.9 ppm). However, the sharp signals exhibit well-resolved J-coupling. This observation can be rationalized with a magnetic, energetically isolated ground state that results from excited state admixture via spin-orbit coupling. Contributions from temperature independent paramagnetism (TIP) are well documented for ³⁰row complexes with d⁶ (octahedral Re⁶, Os⁶) and d⁵ configuration (square-planar, Os⁵). Single-crystal X-ray diffraction confirmed meridional coordination of the neutral diposphinoamine ligand (Figure 1a).
Complex 4 was investigated by cyclic voltammetry (CV) under argon. Quasireversible, scan rate dependent reduction is observed at -1.84 V suggesting chloride loss on the CV timescale. Chemical reduction with 1 equiv. CoCl2 in tetrahydrofuran (THF) under an N2 atmosphere (Scheme 2) gives an intensely blue product (5) in around 60% isolated yield. Complex 5 exhibits sharp and strongly shifted NMR signals suggesting an even-electron compound with a TIP contribution. Symmetric N2 binding was confirmed by a single signal in the 15N[1H]-NMR spectrum and a band at 1733 cm⁻¹ (15N-N2: 1675 cm⁻¹) in the resonance Raman spectrum. Two doublets in the 31P[1H]-NMR spectrum (δp = -370.6/-380.4 ppm) with mutual trans-coupling (Jpp = 237 Hz), eight 1H NMR signals assignable to CH₃-groups and one amine NH-signal are in agreement with an N2 bridged, dinuclear compound with C2 symmetry, in analogy to the molecular structure was confirmed by single crystal X-ray diffraction (Figure 1b). The asymmetric unit features two octahedrally coordinated, linearly N2-bridged Re ions. The N-N bond length (1.169(5) Å) is slightly longer than the C2-N bond (1.202(10) Å), indicating moderate N2 activation in agreement with the Raman data. In contrast to 2, the pincer nitrogen atoms are located trans to the N2 bridge. The two (ReCl3(HPNPPr2)) fragments are twisted with respect to each other by 75.5° giving rise to idealized C2 symmetry. Unlike 2, complex 5 shows remarkable thermal stability. No decomposition was observed over several days, even upon heating to 60 °C in THF. Thermal splitting was therefore examined computationally. Density functional theory (DFT) nicely reproduced the geometry of 5. In analogy to 2, an electronic triplet configuration was obtained as the ground state. While any spin-orbit coupling effects likely present in this dimer are insufficiently described by DFT, the high kinetic barrier (ΔG°298 = 41.8 kcal mol⁻¹) computed for thermal cleavage of 5 into the trans-dichloro nitride [Re(N(trans-Cl2)(HPNPPr2))] (6 trans) is in accordance with experiment and contrasts with cleavage of 2 (ΔG°298 = 26.9 kcal mol⁻¹). Interestingly, splitting of 5 into 6 trans is almost thermoneutral (ΔG°298 = 2.2 kcal mol⁻¹), as compared with strongly exergonic splitting of 2 into square-pyramidal 3 (ΔG°298 = -40.3 kcal mol⁻¹). The less favorable thermochemistry for six-coordinate 6 trans is attributed to the nitride trans-influence as expressed by a long Re-N-Re bond (2.53 Å) and distorted pincer binding. Accordingly, relaxation to isomeric [Re(N)(cis-Cl2)(HPNPPr2)] (6) is strongly favored (ΔG°298 = -11.0 kcal mol⁻¹ per molecule). The striking thermal stability of 5 vs. 2 can be rationalized by qualitative molecular orbital (MO) considerations. Thermal cleavage of a linearly N2 bridged M2N2 core proceeds via electron transfer into an MO with M−N−N−M σ−σ+−σ-character (Scheme 1) within a zig-zag shaped transition state (TS). Ligands trans to the N2-bridge raise this MO in energy, thereby disfavoring N2-splitting. A classic case for such geometry controlled reactivity might be the trisnitriles [[RArN]=Mo2(N2)] 20 which split into nitriles in contrast to analogous triamidoamine complexes. 3a To overcome kinetically hindered N2-splitting we sought to populate N-N antibonding MOs by electronic excitation. Photolysis of 5 in THF with a Xe arc lamp (λ > 305 nm) led to gradual color change to yellow over 2 h. 31P and 1H NMR spectra revealed the formation of cis-dichloro nitride complex 6 (Scheme 2) in 95 % spectroscopic yield upon comparison with an original sample. Use of 15N-labelled 5 confirmed photolytic splitting of the N2 ligand by 15N[1H] NMR spectroscopy. A quantum yield below 1% was estimated by actinometry. The configuration of 6 requires isomerization before or after N2-splitting. Photolysis was therefore carried out in the presence of NHex2Cl (0-500 equiv.). The independence of the reaction rate excludes isomerization by chloride (photo)dissociation prior to or as the rate determining step.
Complex 5 exhibits two maxima above 300 nm in the UV-vis spectrum ($\lambda_{\text{max}} = 336$, $\varepsilon = 38027 \text{ M}^{-1}\text{cm}^{-1}$; 578 nm, $\varepsilon = 12294 \text{ M}^{-1}\text{cm}^{-1}$; Figure 2a). Wavelength selective photolysis indicates that the productive excitations for N$_2$-splitting are centered at a shoulder around 400 nm allowing for the use of a violet LED (390 nm) with the same yield.\textsuperscript{16} TD-DFT reasonably well reproduced the electronic absorption spectrum (Figure 2a) with almost identical results for the open-shell singlet and triplet electronic structure configurations.\textsuperscript{16} Computational evaluation of the highest intensity, yet unproductive absorptions indicate dominant contributions from transitions within the Re=N=N-Re manifold, such as ($\pi^*-$-$\pi^*$)-( $\pi^*-$-$\pi^*$) (calc: 278 nm, 4.46 eV) and ($\pi^*-$-$\pi^*$)-( $\pi^*-$-$\pi^*$) (calc: 460 nm, 2.70 eV). In contrast, two transitions in the productive spectral region (Figure 2a, inset) exhibit mainly (5$d^7$)-( $\pi^*-$-$\pi^*$) character (Figure 2b), implying considerably stronger Re=N$_2$ charge transfer. The weak oscillator strengths ($f_{\text{osc}} \leq 0.02$) are in line with the low quantum yield obtained for photolytic splitting of the N$_2$ bridge.

While the excited state dynamics are not known at this point, the Re=N$_2$ MLCT character of these states is consistent with N$_2$ cleavage. Occupation of Re=N=N-Re ($\pi^*-$-$\pi^*$)-MOs weakens N=N $\pi$-bonding and increases the flexibility of the ReN$_2$-core, which could facilitate cleavage via a zig-zag TS. At the same time, charge separation should enhance Re=N bonding. The spectral features of 5 resemble the visible region of [CP*(depf)Mo(C$_{2}$H$_{4}$N$_2$-$\mu^2$-$\mu^2$-N$_2$)] ($\text{depf} = 1,1'$-bis(diphenylphosphino)ferrocene) which also splits into nitrides upon photolysis at 580 $> \lambda > 400$ nm.\textsuperscript{22a}

In the solid state (Figure 1c),\textsuperscript{16} 6 retains octahedral coordination with one significantly elongated Re–Cl bond (2.4309(7) Å vs. 2.6712(7) Å), reflecting the nitride trans-influence. In turn, the Re≡N bond (1.669(2)Å) is longer compared to five-coordinate [Re(Re)(Cl($\text{H}$PN$^\text{m}$))Cl (1.642(4) Å).\textsuperscript{12a} Addition of NaBAR$_{3}$ (BAR$_{3}$ = tetrakis-[3,5-(trifluoromethyl)phenyl]borate) to 6 in THF results in two new 31P NMR signals that are assigned to two isomers of [Re(N)(Cl($\text{H}$PN$^\text{m}$))]\textsuperscript{16} In turn, 6 is fully restored upon addition of NH$_3$Cl. Hence, six-coordinate [Re(N)(Cl($\text{H}$PN$^\text{m}$))] is the dominant species in solution. Reduced steric shielding facilitates chloride coordination and thereby weakening of metal nitride bonding.

Heating of 6 with PhCO(Cl) (2 equiv.) at 80 °C in 1,4-dioxane over 15 h gives a new rhensium species (7) in 71 % yield (Scheme 2). Paramagnetically shifted and sharp NMR signals support reduction to Re$^0$. Two chemically inequivalent P-atoms ($\delta_p = -1592.6$, -1615.5 ppm; $^2J_{\text{pp}} = 248$ Hz) and the $^1$H and $^13$C NMR signatures are in agreement with pincer oxidation to an imine ligand.\textsuperscript{15} Formation of [Re(Re)(P=NP$^\text{m}$)][(P=NP$^\text{m}$) = N(CH$_2$H$_2$)(CH$_2$H$_2$)(CH$_2$H$_2$)] was confirmed by LIFDI-MS and comparison with an authentic sample.\textsuperscript{16} Crystallographic characterization (Figure 1d) evidences nitride transfer and pincer dehydrogenation.\textsuperscript{16} Two products account for 94 % of the nitride ligand, i.e. benzamide (30 %), and benzonitride (64 % with equimolar benzoic acid), respectively. PhCN and PhCO$_2$H are products from the reaction of benzamide with benzyol chloride,\textsuperscript{23} supporting that PhCO(O)NH$_2$ is the immediate product of nitride benzylation and 2e$^-$/2H$^+$-transfer from the pincer ligand. Related reactivity was observed for [Ru(N($\text{H}$PN$^\text{m}$))], which gives [Ru(NH$_3$)$_2$(O)(CPh)(P=NP$^\text{m}$)] upon reaction with para-methoxy benzoic acid.\textsuperscript{24} However, in that case azide was used as nitrogen source rather than N$_2$.

Next, re-reduction of the imine pincer ligand was examined to evaluate the amine/imine redox couple as cooperating ligand reservoir for 2e$^-$/2H$^+$ proton coupled electron transfer (PCET). 7 does not react with H$_2$ under thermal or photocatalytic conditions. However, chemical reduction is possible with stepwise addition of Li[HBEts] and diphenylammonium chloride (Scheme 2). On this route, 4 is obtained in 62 % spectroscopic yield due to the formation of rhensium hydrides as byproducts, which reform 7 upon hydrolytic quenching.

We therefore turned to electrochemical regeneration of 4. The CV of 7 exhibits quasi-reversible reduction at -1.70 V. Titrating in benzoic acid (0-15 eq.; Figure 3a) results in a pronounced increase of the cathodic current and buildup of a second, quasi-reversible reductive feature at -1.84 V, which is assigned to parent 4. The strong increase of the first wave is indicative of multielectron reduction in the presence of acid. In fact, the viable intermediate [Re(Re)(P=NP$^\text{m}$)] (8) exhibits a quasi-reversible reduction at -1.16 V,\textsuperscript{16} confirming a potential inversion after the first e$^-$/H$^+$-transfer to 7. The electrochemical data in the presence of acid is therefore rationalized with two subsequent PCET steps at around -1.7 V that regenerate 4.

Controlled potential electrolysis (CPE) of 7 at the half peak potential of the first reduction feature (E = -1.65 V) did not give appreciable amounts of 4 in the presence of benzoic acid (10 eq). 2,6-Dichlorophenol (DCP) was therefore employed, which exhibits about same pKa as benzoic acid (pKa$_{\text{THF}}$ = 25.1)$^{25}$ yet a conjugate base that is less prone to metal coordination. In fact,
bulk electrolysis of 7 at $E = -1.65$ V in the presence of DCP (10 eq.) for 7 h results in full consumption of 7 and formation of 4 as the only electroactive species. $^1$H NMR spectroscopy confirmed a yield of 99% and coulometry the transfer of 1.96 e\textsuperscript{-} per 7.

The quantitative regeneration of 4 motivated the examination of \textit{in situ} electrochemical N\textsubscript{2} activation. In a CPE experiment, 7 was electrolyzed with DCP (10 eq.) under N\textsubscript{2} for 8 h at $E = -1.65$ V, followed by 5 h at $E = -1.85$ V, i.e. the half potential of the reduction of 4 (Figure 3b). Overall, 3.32 e\textsuperscript{-} per Re atom were transferred. Formation of complex 5 during the second electrolysis step was indicated by the deep blue color and confirmed by $^1$H and $^{31}$P($^1$H) NMR spectroscopy. UV/Vis spectroscopic quantification gave a yield around 70% (Scheme 3). In situ photolysis (390 nm LED) of this mixture revealed N\textsubscript{2} splitting to nitride 6 as the only detectable product by $^{31}$P($^1$H) and $^1$H NMR spectroscopy. However, the yield in 6 dropped to 14% with respect to parent 7, i.e. considerably lower compared to photolysis of isolated 5.

In summary, a three-step cycle for the generation of benzamide/benzonitrile from N\textsubscript{2} in overall 61% yield (with respect to N) was established that relies on electrochemical N\textsubscript{2} activation, photochemical splitting into nitrides and thermal nitrogen transfer (Scheme 3). This model reveals some basic principles in comparison to our previously reported system. Use of a sterically less encumbered pincer ligand stabilizes higher coordination numbers. In consequence, thermal N\textsubscript{2} splitting becomes less favorable which can be overcome by photolysis with visible light. Based on the experimental and theoretical data, the photochemical reactivity is associated with the population of a dissociative state with Re–N\textsubscript{2} MLCT character. The higher coordination number weakens nitride bonding, thus enabling the use of a weaker electrophile than alkyl triflates. Importantly, the cooperating pincer ligand serves as a reservoir for nitrogen hydrogenolysis upon 2e\textsuperscript{-}/2H\textsuperscript{+}-PCET and electrochemical re-hydrogenation. Our results demonstrate how metal-ligand cooperativity and photo- and electrochemical approaches can facilitate the design of platforms for N\textsubscript{2} fixation.

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[16] For synthetic, spectroscopic, computational and crystallographic details see Electronic Supporting Information (ESI).


[18] All electrochemical data is reported vs. FeCp₂/FeCp₂⁺.


Cooperative Fixation: $N_2$ fixation to benzonitrile and benzamide is reported within a three-step cycle that utilizes electrochemical reduction, photochemical $N_2$ splitting and thermal $N$-transfer with benzoylchloride. Product formation is enabled by cooperativity of the functional pincer ligand, which serves as a $2H^+/2e^-$ reservoir.