Molecular Electrochemical Reductive Splitting of Dinitrogen with a Molybdenum Complex

Lydia Merakeb, Soukaina Bennaamane, Eric Clot, Nicolas Mézailles and Marc Robert

Université de Paris, Laboratoire d’Electrochimie Moléculaire, CNRS, F-75006 Paris, France
Laboratoire Hétérochimie Fondamentale et Appliquée – UMR 5069, Université Toulouse III – Paul Sabatier, 118, route de Narbonne, Bât. 2R1, 31062, Toulouse, France
ICGM Univ Montpellier, CNRS, ENSCM, Montpellier, France
Institut Universitaire de France (IUF), F-75005, Paris, France

KEYWORDS. Dinitrogen reductive splitting, electrochemical reduction of N₂, molybdenum complex, nitride Mo complex

ABSTRACT: Nitrogen reduction in mild conditions (i.e. room temperature and atmospheric pressure) and using a non-fossil source of hydrogen remains a high chemistry challenge. Molecular metal complexes, notably Mo based, have recently shown to be active for such nitrogen fixation. In this work, we report about the electrochemical N₂ splitting with MoV triphosphino complex ((PP)MoI), at room temperature and a moderately negative potential. A MoIV nitride species was generated, which was confirmed by electrochemistry and NMR studies. The reaction goes through the bi-electronic reduction of the starting Mo species, coordination of an N₂ molecule, and further splitting to a MoV nitride complex. Preliminary DFT investigation supports the intermediacy of a bridging MoN₂Mo nitride dimer evolving to the Mo nitride via a low energy transition state. This example joins a short list of molecular electrochemical complexes for N₂ reductive cleavage. It opens a door to molecular electrochemical PCET conversion studies of N₂ to NH₃.

INTRODUCTION

Nitrogen is one of the essential elements for life.¹ Although abundant, it is nonetheless a limiting nutrient in agriculture, and crops growth is dependent on its availability.² It is not only essential to the global economy as a fertilizer but also as the feedstock for industrial production of all N containing derivatives. Moreover, it has been identified as an alternative fuel as well as an energy storage molecule.³⁴ Today, ca. 200 million tons of NH₃ are produced yearly, exclusively by the Haber-Bosch (H-B) process (equation (1)).

\[
\text{N}_2 + 3 \text{H}_2 \rightleftharpoons 2 \text{NH}_3 \quad \Delta G^\circ = 32.9 \text{ kJ mol}^{-1}
\]  

Although thermodynamically favorable, the reaction is carried out at high temperature (300 - 500 °C) and high pressure (200 - 300 bars) conditions over heterogeneous Fe containing catalysts, using ca. 1-2% of the global energy production. It generates half a Gigaton of CO₂/year (ca. ½ of the CO₂ generated is due to H₂ production by CH₄ reforming), a giant carbon footprint.

Nature, by the means of microorganisms, transforms N₂ into NH₃ under ambient conditions, via multiple proton/electron transfers.⁵ Nitrogenase enzymes, responsible for this “N₂ fixation” reaction (equation (2)), contain an organometallic active site, a cluster “MoFe/S₆C-homocitrate” which can be viewed as a homogeneous catalyst. The fact that this catalyst involves a coordination/activation of N₂ at one/several metal centers has spurred “organometallic chemists” to search for homogeneous catalysts able to achieve the N₂-to-NH₃ reaction efficiently.

\[
\text{N}_2 + 8 \text{H}^+ + 8 \text{e}^- + 16 \text{ATP} \rightleftharpoons 2 \text{NH}_3 + \text{H}_2 + 16 \text{ADP} + 16 \text{P}_i
\]  

Transition metal complexes based on Mo⁶⁻¹² and Fe¹³⁻¹⁶, mainly, but also rhodium¹⁷⁻¹⁹, vanadium²⁰, chromium²¹⁻²³, cobalt²⁴⁻²⁵ and titanium²⁶ have been investigated. However, a major hurdle toward such chemistry lies in the kinetically favorable proton reduction into H₂. Thus, despite several decades of active research, only a handful of catalysts are known for this reaction. Regarding electrochemical nitrogen splitting with molecular complexes,²⁷⁻²⁹ an initial example was identified by Picket and Talarmin with a W complex,³⁰ and in recent years, work has been performed on Ti³¹ and Al³² based complexes as well. The first and only so far example of electrochemically driven N₂-to-NH₃ catalytic conversion with a well-defined molecular species was reported in 2016. P₈⁵Fe(N₂)²⁻ (P₈⁵ = tris(o-diisopropylphosphinophenyl)borane) converts N₂ to NH₃ in Et₂O containing 0.1 M Na[BAr₄²⁻] as a supporting electrolyte (BAr₄²⁻ = tetrakis(3,5-bis(trifluoromethyl)phenyl)borate) at – 45 °C (Scheme 1a). With 50 eq. of H[BAr₄²⁻] as a proton source and upon applying a potential of – 2.3 V vs. Fe/Fe⁺, 2.2 eq. of ammonia were produced with 25 % Faradaic efficiency over 16.5 h.³³
More recently, electrochemical reduction of N₂ to hydrazine was shown on a tri-nickel based molecular catalyst. In THF (0.1 M Bu₄NPF₆), N₂ was converted into N₂H₄ at -2.35 V vs. FeC/Fc⁺ with 8 to 12% Faradaic efficiency, in the presence of 0.1 M phenol as a proton source (Scheme 1b).³⁴

Electrochemical N₂ activation through splitting of the triple bond into metal nitrides has been recently the focus of several studies. Schneider, Siewert and coworkers have used a Re⁺⁺ complex for electrochemical conversion into Re⁺ nitride in 58% yield at E = -1.9 V vs. FeC/Fc⁺ in presence of dinitrogen (Scheme 1c). Using a rhodium complex with a similar but more conjugated pincer ligand led to N₂ splitting at a less cathodic potential, however at the expense of a lower yield (Scheme 1d).³⁵ Remarkably, Masuda et al. showed that splitting of dinitrogen into terminal nitrides is also possible by chemical and electrochemical oxidation of a N₂-bound Mo⁺⁺ complex (Scheme 1e).³⁶ Still regarding molybdenum, we have shown that the splitting of N₂ could be achieved through the two electron reduction of a (PPP)MoCl₃ complex (Scheme 2a) in the presence of NaI, and subsequently studied the functionalization of the nitride complex by silanes and boranes.¹⁰ These experimental studies, coupled to DFT calculations demonstrated the facile transformation of the nitride to imido (boryl and silyl) and to amido derivatives, which are related to the key intermediates toward complete reduction to ammonia.³⁷ In this work, we achieved the electrochemical splitting of dinitrogen with a Mo³⁺ trim-phosphino complex (PPP)MoI₃ (Mo₂, Scheme 2b) that leads to the formation of the corresponding Mo–nitride complex, at room temperature and moderate potential.

**RESULTS AND DISCUSSION**

The cyclic voltammetry of Mo₂ under Ar shows four distinct features (Figure 1). A first reduction wave was observed at -1.15 V vs. SCE. The thin wave is indicative of a fast charge transfer and comparison to reversible couples of similar size shows that reduction is mono-electronic. It could be assigned to Mo³⁺/Mo⁺⁺ couple. Irreversibility of this wave is due to a chemical step following the initial electron transfer, most likely iodide ligand dissociation. Indeed, oxidation waves at 0.33 and -1.15 V vs. SCE are indicative of a chemical splitting of dinitrogen with a Mo³⁺ trim-phosphino complex (PPP)MoI₃ (Mo₂, Scheme 2b) that leads to the formation of the corresponding Mo–nitride complex, at room temperature and moderate potential.

**Figure 1.** CVs of Mo₂ (0.58 mM) in THF + 0.1 M Bu₄N[BarF₄] at a glassy carbon electrode, under Ar (black) and N₂ (red), at v = 100 mV/s and room temperature (RT).

**Figure 2.** CVs of Mo₂ (1 mM) in THF + 0.1 M Bu₄N[BarF₄] under N₂ atm, at v = 500 mV/s and RT. The starting potential was held for 15 s (black, green, orange and red). Blue trace: CV of Mo₃ (1.12 mM) in THF + 0.1 M Bu₄N[BarF₄] under N₂ atm, at v = 500 mV/s, RT.
0.61 V vs. SCE are observed, corresponding to I\(^{-}\) oxidation to I\(^{2-}\) and I\(^{3-}\) respectively. A second irreversible wave is located at -1.8 V vs. SCE (Figure 1, pc3). It may be noted that great care should be taken in the choice of supporting electrolyte anion. The classical hexafluorophosphate (PF\(_6\)) does coordinate to Mo, preventing any further reactivity with N\(_2\). The bulky tetrakis(3,5-bis(trifluoromethyl)phenylborate (BAr\(^{+}\)) is non-coordinating and should be preferred.

Under an atmosphere of N\(_2\), the cyclic voltammetry of Mo2 shows additional features. A new reduction peak appears at -1.40 V vs. SCE (Figure 1, pc2), along with a related oxidation peak at -0.26 V vs. SCE (Figure 1, pa). This later peak is only observed when the potential is scanned until pc2, thus it corresponds to the oxidation of a species formed at this later wave. Both waves are diffusion limited. The irreversible pc3 wave is not affected by the presence of N\(_2\) (this later reduction will not be further investigated.). On the contrary, intensity of the first reduction peak (Figure 1, pc1) increases slightly and the peak potential is positively shifted, suggesting possible coordination to dinitrogen. The initial electrochemical behavior could be restored upon Ar bubbling.

To identify the species formed at pc2 and oxidized at pa, CVs of an authentic sample of Mo3 were recorded in the same conditions. Oxidation of this complex occurs at the exact same potential as the species formed at pc2, as shown in Figure 2. This indicates the likely formation of the nitride complex Mo3 at pc2 reductive wave.

![Figure 3](image-url)

**Figure 3.** a) CV at 1 mm GC disk of the electrolyte solution after CPE (black) and CV at 3 mm GC disk of Mo3 (1.12 mM) in THF +0.1 M Bu4N[Bar\(^{+}\)] under N\(_2\) atm., at v = 100 mV/s, RT. b) \(^{31}\)P\([\text{I}^\text{H}]\) NMR of the same electrolyte solution. c) \(^{31}\)P\([\text{I}^\text{H}]\) NMR of complex Mo3 in THF d-8.

To further investigate and demonstrate N\(_2\) reductive splitting, a controlled potential electrolysis (CPE) of a THF solution (+0.1 M Bu4N[Bar\(^{+}\)]) containing Mo2 and saturated with N\(_2\) was performed at -1.4 V vs. SCE, i.e. at pc2 peak potential. After a few minutes, coloration of the electrolyte solution changed from orange to greenish. After 35 min, a charge of 1.03 C was exchanged, corresponding to 1.24 e\(^{-}\) per Mo centre (see Supporting Information for details). At the end of the electrolysis, cyclic voltammetry of the electrolyte solution was performed (Figure 3). The oxidation waves observed at ca. 0.5 and 0.75 V vs. SCE correspond to oxidation of iodide ions to I\(^{2-}\) and I\(^{3-}\) and the reduction wave at ca. -1.75 V vs. SCE corresponds to the same process characterized by the wave pc3 in Figure 1. Most importantly, a new reversible wave is observed at -0.31 V vs. SCE. This wave corresponds to the reversible oxidation of Mo\(^{IV}\) nitride complex Mo3, generated electrochemically during the CPE. Indeed, comparison with CVs of an authentic sample of Mo3 yields a reversible wave at the exact same potential (Figure 3a).

\(^{31}\)P-NMR provided further characterization of the nitride complex. After the CPE, the electrolyte solution was concentrated in a J-Young NMR tube, and a small volume of deuterated THF was added. Figure 3b shows two singlet peaks at 69 and 121 ppm respectively, which are characteristic of complex Mo3 as deduced from comparison with the \(^{31}\)P\([\text{H}]\) spectrum of an authentic sample Mo3. The two additional singlet peaks observed at 53 and 98 ppm also correspond to a high oxidation state complex and are nearly identical to the spectral signature of the oxoo complex [(PPP)Mo=O(OCl)\(_2\)] (53.2 and 98.7 ppm respectively). This later species is likely formed during electrolysis. When the same electrolysis experiment was repeated under an Ar atmosphere, neither the reversible wave at -0.31 V vs. SCE in cyclic voltammetry, nor the signals at 69 and 121 ppm in \(^{31}\)P\([\text{H}]\) NMR were observed. On the other hand, the oxo complex is formed efficiently under these conditions (see ESI). Understanding the precise mechanism of formation of this complex is beyond the scope of the present work, and is currently studied in our laboratories. Nonetheless, preliminary experiments indicate that both O\(_2\) and H\(_2\)O can be ruled out as the source of “oxygen” for the oxo species.
Scheme 4. DFT mechanistic insights into reductive N₂ splitting with Mo₃.

Combined CV and ³¹P NMR data converge to the robust conclusion that Mo₃ is obtained from Mo₂ and dinitrogen upon electrochemical reduction. The yield of the reaction could be derived from cyclic voltammetry by comparing the intensity of the observed reversible wave at –0.31 V after CPE with a CV recorded in the same electrolyte conditions, before CPE. The yield was found to be ca. 30%. This is in agreement with the charge passed during the electrolysis (1.24 e⁻ per Mo). Full conversion of Mo₂ to Mo₃ would require 2e⁻ per Mo centre, and it was already apparent it was not the case upon inspection of cyclic voltammetry data (Figures 1 and 2). Indeed, the intensity of the pc2 wave, at which Mo₃ is produced, is significantly smaller than that of the one electron pc1 wave.

Preliminary DFT calculations were carried out to get insights into the reaction mechanism (Scheme 4, see ESI for details). Relative energies between Mo complexes with different redox states were estimated using Cp⁹Co as a one electron reducing agent. The lowest potential energy surface (PES) (lowest energy complexes with corresponding spin state) is presented. Complexes in different spin states have been computed and are mentioned when relevant. Calculations start with the [(PPP)MoI₂] complex ^4A, a quartet of spin. One electron reduction leads to the anionic complex ^3B (quintet of spin) in a mildly exergonic process (–3.4 kcal mol⁻¹). Interestingly, one of the Mo-I apical bonds is strongly elongated in ^3B (5.38 Å), and thus the unsaturated complex ^5C [(PPP)MoI₂] lies close in energy (–0.6 kcal mol⁻¹ with respect to ^3B). Calculations could not locate a local extremum associated to N₂ coordination to ^5C. As observed in the tri-iodide structure, octahedral geometry is not favored with a quintet spin state (Mo…N₂ distance of 4.45 Å). However, the complex ^5C with a triplet spin state lies slightly above ^5C (–2.8 and –4.0 kcal mol⁻¹ resp.) and coordination of N₂ to this complex is a kinetically facile step. Indeed, the transition state TS(^1C–^2D) is located only 15.9 kcal mol⁻¹ higher than ^3C. Dinitrogen Mo(II) complex ^3D is only 1.6 kcal mol⁻¹ higher than ^5C, and these two complexes are therefore in equilibrium. Splitting of N₂ was evaluated at the Mo⁹ redox state. Reaction between ^3C and ^3D to form a dimer ^3C–^3D (triplet spin state) or ^1C–^3D (singlet spin state) is computed to be endergonic with respect to ^3C and ^3D (ΔG = 9.6 kcal mol⁻¹, ^3C–^3D; ΔG = 4.9 kcal mol⁻¹, ^1C–^3D). Moreover, calculations of the transition states for N₂ cleavage for both dimers resulted in very high activation energies (ΔG° = 52.0 kcal mol⁻¹, from ^1C–^3D; ΔG° = 37.3 kcal mol⁻¹, from ^3C–^3D), and such pathway can be discarded.

One electron reduction of ^3C and ^3D is computed to be endergic by ΔG = 5.4 kcal mol⁻¹ and ΔG = 6.1 kcal mol⁻¹, respectively. The reduced species are more stable as doublet spin states and iodide dissociation yields the mono-iodo complex [(PPP)MoI] (^6E) and ^7F from coordination to N₂. The combined reduction / iodide dissociation is computed to be endergic in the case of ^3C → ^6E + I (ΔG = 4.3 kcal mol⁻¹), whereas it is slightly exergic in the case of ^3D → ^7F + I (ΔG = –0.4 kcal mol⁻¹). Contrary to the situation encountered with Mo⁹, the singlet–N₂ complex ^7F is computed to be more stable than the unsaturated Mo¹ complex ^6E by ΔG = –3.1 kcal mol⁻¹. As a result of this additional stability, the N₂ coordination to ^6E is computed to be favorable (ΔG° = 8.4 kcal mol⁻¹) to yield ^7F. Formation of dimer species between ^6E and ^7F, either as a singlet (^6E–^7F) or as a triplet (^6E–^7F), is computed to be exergonic (ΔG = –15.2 kcal mol⁻¹, ^6E–^7F; ΔG° = –15.4 kcal mol⁻¹, ^6E–^7F). Despite being close in energy, the N₂ cleavage is operative only from the singlet dimer ^6E–^7F with an activation barrier of ΔG° = 21.0 kcal mol⁻¹, whereas the process is computed to be highly unfavorable from the triplet dimer (ΔG° = 65.0 kcal mol⁻¹). The singlet transition
state exhibits an expected zig-zag geometry with a highly elongated N-N bond (1.631 Å vs. 1.192 Å in $^{1}$E-F) and shorter Mo-N contacts (1.689 Å and 1.693 Å vs. 1.868 Å and 1.871 Å in $^{1}$E-F). The overall transformation $^{3}$E + $^{4}$F $\rightarrow$ $^{2}$G, where $^{1}$G is the Mo$^{V}$-nitrido (Mo$^{3}$) final product resulting from N$_{2}$ cleavage is computed to be strongly exoergic ($\Delta G = -55.3$ kcal mol$^{-1}$).

Overall, DFT studies support the electrochemical mechanism for N$_{2}$ splitting given in Scheme 3. It involves spontaneous iodide dissociation upon one-electron reduction to form the Mo$^{6}$ di-iodide intermediate in equilibrium with the N$_{2}$-coordinated Mo$^{5}$ complex. Mo$^{6}$ N$_{2}$-dimer formation is thermodynamically accessible but N$_{2}$ cleavage is kinetically prohibited. A second reduction at more negative potential results in further iodide dissociation and easy N$_{2}$ coordination to Mo$^{5}$, resulting in thermodynamically easy formation of N$_{2}$-dimer species from which splitting to Mo$^{V}$ nitride species is kinetically accessible.

In summary, we have demonstrated the electrochemical splitting of dinitrogen at a Mo molecular complex in ambient conditions, and characterized the formation of the corresponding Mo-nitride complex. To the best of our knowledge, this is the first electrochemical reduction driven splitting of N$_{2}$ with molybdenum species. Such nitride complexes are ideal starting point not only for the chemical functionalization but also and most importantly for electrochemical studies aiming at generating ammonia upon successive proton-coupled electron transfers (PCET process), one step closer from molecular electrochemical catalysis of N$_{2}$ reduction with metal complexes.

ASSOCIATED CONTENT
Materials and methods, supplementary CV data and NMR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author
* Email: mezailles@chimie.ups-tlse.fr
* Email: robert@u-paris.fr

ORCID
Lydia Merakeb: 0000-0002-5834-8144
Soukaina Bennaamane: 0000-0002-5358-0585
Eric Clot: 0000-0001-8332-5545
Nicolas Mézaillès: 0000-0001-8735-6285
Marc Robert: 0000-0001-7042-4106

Author Contributions
The manuscript was written through contributions of all authors. * All authors have given approval to the final version of the manuscript. ‡ These authors contributed equally.

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Notes
The authors declare no competing financial interest.

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Supporting information

Molecular Electrochemical Reductive Splitting of Dinitrogen with a Molybdenum Complex

Lydia Merakeb\textsuperscript{a}, Soukaina Bennaamane\textsuperscript{b}, Eric Clot,\textsuperscript{c} Nicolas Mézailles\textsuperscript{b}\textsuperscript{*} and Marc Robert\textsuperscript{a,d}\textsuperscript{*}

\textsuperscript{a}Université de Paris, Laboratoire d’Electrochimie Moléculaire, CNRS, F-75006 Paris, France
\textsuperscript{b}Laboratoire Hétérochimie Fondamentale et Appliquée – UMR 5069, Université Toulouse III – Paul Sabatier, 118, route de Narbonne, Bât. 2R1, 31062, Toulouse, France
\textsuperscript{c}ICGM, Univ Montpellier, CNRS, ENSCM, Montpellier, France
\textsuperscript{d}Institut Universitaire de France (IUF), F-75005, Paris, France
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1. Materials and Methods

All experiments were performed under either Ar or N₂ atmosphere using a glove-box and standard Schlenk-line techniques. THF, Et₂O and DCM (HPLC grade) were purchased from VWR. THF and Et₂O were distilled and further dried over 3 Å molecular sieves for at least 48 h before use. DCM was distilled over CaH₂. Deuterated solvents were purchased from Eurisotop and THF-d₈ was dried over 3 Å molecular sieves. ACN (99.9 % extra dry over molecular sieves) was purchased from Acros Organics and was further dried over an activated alumina column (neutral Brockman Activity I) before use. Ar and N₂ (alphagaz 2, Air Liquide) were further dried over CaCl₂ when not used in the glove-box.

2. Analytical methods

Spectroscopy. NMR spectra were recorded on a Bruker 400 MHz spectrometer. ¹H spectra were calibrated using the residual solvent signals and ³¹P spectra were reported relative to phosphoric acid as external standard.

Electrochemistry. All electrochemical data were collected on a Metrohm Autolab instrument. For CV measurements, the working electrode was a homemade 3 mm glassy carbon disk (Tokai) sealed in glass. It was carefully polished using diamond paste (1 µm), rinsed with acetone and dried before each use. The platinum thread counter-electrode was flame annealed before use. The reference electrode was a non-aqueous Ag/AgNO₃ quasi-reference electrode, composed of a flame annealed Ag thread in acetonitrile containing 0.1 M Bu₄NPF₆ and 0.01 M AgNO₃ in a Vycor tip terminated bridge. It was separated from the working electrode compartment by a fritted glass bridge. The exact potential of this reference vs. SCE was measured using a multimeter before and after each experiments. Potentials are referred to SCE in all figures. Ohmic drop was compensated using the positive feedback implemented in the instrument.

In Controlled Potential Electrolyses (CPEs), the working electrode was a polished 1 cm² carbon plate, whose surface was defined using Teflon tape. The counter-electrode was a flame annealed Pt grid in THF containing 0.1 M Bu₄N[BArF₄] and 100 mM ferrocene as a sacrificial oxidation substrate and the reference electrode was a non-aqueous Ag/AgNO₃ electrode. Both CE and RE were separated from the main compartment by two fritted glass bridges. The electrolyte solution was kept under forced convection with a magnetic stirrer.

3. Syntheses

Synthesis of Mo₂ and Mo₃ Mo₃ was synthesized following a reported procedure.¹ Mo₂ was prepared as follows: [MoI₃(THF)₃]²⁺ (100 mg, 0.146 mmol) was dissolved in 20 mL THF and stirred for a few minutes. Then 0.86 mL of a solution of P³¹P₂Cy in THF (0.17M, 0.146 mmol) was added and the mixture was stirred for 24 hours at room temperature. The solution turns from dark red to orange and Mo₂ starts precipitating as an orange powder. The solvent is then evacuated in vacuo and the powder is washed twice with pentane and dried under vacuum. Isolated yield: 134 mg, 89%. MALDI-TOF: calculated for C₃₄H₅₇I₂MoP₃: 910.08; found: 910.11.
Synthesis of Bu₄N[BAr₄F₄] and Na[BAr₄F₄]

Sodium n-tetrabutylammonium tetrakis(3,5-bis(trifluoromethyl)phenyl) borate was synthesized by metathesis of sodium tetrakis(3,5-bis(trifluoromethyl)phenyl) borate with n-tetrabutylammonium chloride. Na[BAr₄F₄] was dissolved in a minimal amount of milliQ water, to which methanol was added till the salt was completely dissolved. 1.05 eq. of Bu₄Cl were also dissolved in a minimal amount of water, and added dropwise, under vigorous stirring. Bu₄N[BAr₄F₄] formed immediately as a white precipitate. It was then collected by filtration, rinsed several times with milliQ water and dried at 80 °C under vacuum for 48 hours.

Na[BAr₄F₄] was synthesized following a reported procedure.²

Synthesis and characterization of the oxo complex

[Mo(PPh₂P₂Cy)(Cl₂)(O)] ([PPPMo=O(Cl)_2])

[Mo(PPh₂P₂Cy)(Cl)](THF)_2 (194.0 mg, 0.215 mmol) and NaI (64.6 mg, 0.431 mmol) were dissolved in “wet” THF (10 mL, containing ca 500 ppm H₂O) in a Schlenk flask. The mixture was stirred for 10 min at room temperature under 1 atm N₂. Then Sodium amalgam was added (0.431mmol). After stirring 5 hours, the solution had turned from yellow to green. An aliquot was taken and analyzed in ³¹P NMR, revealing two products, the known nitrido complex [Mo(PPh₂P₂Cy)NI] as a minor product and a new diamagnetic compound as the major product. The solution mixture was filtered then the solvent was partially evacuated in vacuo. The obtained concentrated solution was placed in vial and a couple of drops of pentane were added slowly, then the vial was kept at -30°C. After 2 days, green crystals were obtained in 70% yield. The X-Ray diffraction analysis showed these crystals to be the oxo complex [Mo(PPh₂P₂Cy)(Cl)_2(O)].

¹H NMR (400 MHz, THF-d₈, 25 ºC) δ:  7.7 (m, 2H, 2 H_o Ph), 7.4 (m, 3H, H_p Ph, 2 H_m Ph), 2.9 – 0.7 (m, 4 CH_Cy, 20 CH₂Cy, 4 P(CH₂)₂P) ppm.

³¹P{¹H} NMR (160 MHz, THF-d₈, 25 ºC) δ: 53.2 (s, 2P), 98.7 (s, 1P) ppm.

MALDI-MS (m/z): 742 [M]+

Figure S1. ³¹P{¹H} NMR spectrum of Mo=O.
Figure S2. $^1$H NMR spectrum of Mo=O.

Table S1. Crystal data and structure refinement for Mo=O.

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<th>Property</th>
<th>Value</th>
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<td><strong>Empirical formula</strong></td>
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<td><strong>Formula weight</strong></td>
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<td><strong>Temperature</strong></td>
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<tr>
<td><strong>Space group</strong></td>
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<tr>
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<tr>
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<tr>
<td>b</td>
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</tr>
<tr>
<td>c</td>
<td>21.1229(9) Å</td>
</tr>
<tr>
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<tr>
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<tr>
<td><strong>Density (calculated)</strong></td>
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<tr>
<td><strong>Absorption coefficient</strong></td>
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**Table S2.** Selected bond lengths [Å] and angles [°] for Mo=O.

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<tr>
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<th>Bond Angles (°)</th>
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<td>Mo(1)-P(2)</td>
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</table>

- Bond Distances
- Bond Angles

- Crystal size: 0.200 x 0.200 x 0.160 mm³
- Theta range for data collection: 3.157 to 33.699°
- Index ranges: -20 ≤ h ≤ 20, -20 ≤ k ≤ 20, -32 ≤ l ≤ 31
- Reflections collected: 156420
- Independent reflections: 14494 [R(int) = 0.0626]
- Completeness to theta = 25.242°: 99.8%
- Absorption correction: Semi-empirical from equivalents
- Max. and min. transmission: 0.7384 and 0.6949
- Refinement method: Full-matrix least-squares on F²
- Data / restraints / parameters: 14494 / 0 / 370
- Goodness-of-fit on F²: 1.052
- Final R indices [I>2sigma(I)]: R1 = 0.0364, wR2 = 0.0724
- R indices (all data): R1 = 0.0678, wR2 = 0.0850
- Extinction coefficient: n/a
- Largest diff. peak and hole: 1.550 and -1.009 e.Å⁻³
4. CV data

**Figure S3.** CVs at 3 mm GC electrode in THF (0.1 M Bu4N[Bar2]) containing 1.2 mM FeTPPCl (black) or 0.58 mM Mo2 (red and green). At $v = 100$ mV/s, under Ar. CVs are normalized for concentration C and number of exchanged electrons n (red trace: n = 1, green trace: n = 2).

**Figure S4.** CVs at 3 mm GC electrode in THF (0.1 M Bu4N[Bar2]) containing 1 mM Mo2. $v = 100$ mV/s, under N2.
5. Controlled Potential Electrolyses

**CPE under N₂**
A controlled potential electrolysis was performed in THF (0.1 M Bu₄N[BarF₄]) containing ca. 2.5 mM Mo₂, at – 1.4 V vs. SCE, *i.e.* at the pc2 peak potential (see main text), in a N₂ filled glovebox. A charge of 1.03 C was passed, corresponding to 1.24 e⁻ per Mo atom. Before and after the electrolysis, CVs were recorded at a 1 mm glassy carbon electrode. After the CPE, the electrolyte solution was placed in J-young terminated NMR tube and concentrated, and NMR spectra were recorded.

![Figure S5](image_url) CVs at 3 mm GC electrode in THF (0.1 M Bu₄N[BarF₄]) containing 2.46 mM Mo₂ under N₂.

![Figure S6](image_url) CA at 1 cm² GC electrode in THF (0.1 M Bu₄N[BarF₄]) containing 2.46 mM Mo₂, under N₂.
**CPE under Ar**

A solution of 2.5 mM Mo2 in THF (0.1 M Bu₄N[BArF₄]) was prepared and placed in the electrochemical cell in the N₂ filled glovebox, then degassed under an Ar stream for more than 30 min. A CPE was performed at –1.4 V vs. SCE. After the electrolysis, 0.37 C were exchanged, corresponding to 1 e⁻/Mo center.

![Graph](image1)

**Figure S7.** CV at 1 mm GC electrode in THF (0.1 M Bu₄N[BArF₄]) containing 1 mM Mo2 before electrolysis. At v = 100 mV/s, under Ar.

![Graph](image2)

**Figure S8.** CA at 1 cm² GC electrode in THF (0.1 M Bu₄N[BArF₄]) containing 1 mM Mo2, under Ar.
6. NMR data

Supporting electrolytes

Figure S10. $^1$H NMR spectrum of Na[Bar$^2$] in CD$_3$CN. * Solvent residual peaks (H$_2$O at 2.16 ppm and CH$_3$CN at 1.94 ppm).
Geometry optimizations were performed using Gaussian 16 (Revision A03) at the PBE0 level of hybrid density functional theory, with inclusion of D3(bj) corrections in the optimization process. The geometries of all located extrema are given as xyz coordinates data in a separate file (Geom.xyz) in the SI. The atoms Co, C, H, N and P were represented by an svp basis set. The Mo and I atoms were represented by Dolg’s pseudo potential and the associated basis set. The solvent (thf) influence was taken into consideration through single-point calculations on the gas-phase optimized geometries with SCRF calculations within the SMD model. For the SCRF calculations, the atoms were treated with a def2-qzvp basis set. All energies reported are Gibbs free energies obtained by summing the SMD energy (including D3 corrections) and the gas phase Gibbs contribution at 298 K and 1atm.
Relative energies (as compared to $^5A$) of all computed structures (Scheme 4, see main text) are given below. Exponent is the spin state.

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8. Abbreviations

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<tr>
<th>Abbreviation</th>
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<tr>
<td>BAr$_4^+$</td>
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9. References


