A series of cationic diazoalkane complexes \([4-\text{RC}_6\text{H}_4\text{C(H)NNM}o(\text{N}[\text{t-Bu}\text{Ar}])_3][\text{AlCl}_4]\), \([1-\text{R}][\text{AlCl}_4]\) (\(\text{R} = \text{NM}e_2, \text{Me}, \text{H}, \text{Br}, \text{CN}, \text{Ar} = 3,5-\text{C}_6\text{H}_2\text{Me}_2\)) have been prepared by treatment of the \(\text{N}_2\)-derived diazenido complex \(\text{Me}_3\text{SiNNM}o(\text{N}[\text{t-Bu}\text{Ar}])_3\) with \(4-\text{RC}_6\text{H}_4\text{CHO}\) and 2 equiv \(\text{AlCl}_3\). The structures of \([1-\text{H}][\text{AlCl}_4]\) and \([1-\text{NM}e_2][\text{AlCl}_4]\) were determined by X-ray crystallography. The \(\text{C}–\text{N}\) and \(\text{N}–\text{N}\) stretching modes were identified by a combined IR and Raman spectroscopy study, and other physical properties are discussed in detail. The electrochemical reduction potential for \([1-\text{R}][\text{AlCl}_4]\) was shown to be linear with the Hammett \(\sigma\) parameter. This reduction process forms the \(\text{C}–\text{C}\) bonded dimer, \(\mu-(4-\text{RC}_6\text{H}_4\text{C(H)NN})_2[\text{Mo(N}[\text{t-Bu}\text{Ar}])_3]\) that was characterized by X-ray crystallography for \(\text{R} = \text{H}\). Possible mechanisms for the formation of this dimer are presented. Both electrochemical investigations and quantum chemical calculations are used to describe the odd-electron complex \(4-\text{RC}_6\text{H}_4\text{C(H)NNM}o(\text{N}[\text{t-Bu}\text{Ar}])_3, 1-\text{R}\), that is an intermediate in formation of \([1-\text{R}]_2\). The \(\text{C}–\text{C}\) bond in \([1-\text{R}]_2\) is redox non-innocent and is broken upon oxidation. This reaction was used to prepare \([1-\text{H}]\text{[A]}\) (\(\text{A} = \text{PF}_6, \text{OTf}\)), and possible uses of this property in charge-storage devices are discussed.

Introduction

A great deal of energy has been expended optimizing methods for the chemical fixation of dinitrogen.1–7 Of these methods, the Haber-Bosch process, that forms \(\text{NH}_3\) from \(\text{N}_2\) and \(\text{H}_2\), is the most well-studied and currently the most widespread process for nitrogen fixation.8 Ammonia produced by this process may be incorporated into organic molecules in a subsequent synthetic step. However, a more synthetically efficient route to nitrogen-containing organic molecules would directly incorporate \(\text{N}_2\) into organic fragments without the intermediacy of ammonia.9–13 Such a synthetic protocol could be designed to avoid the need for \(\text{H}_2\), which currently is formed by steam reforming of \(\text{CH}_4\) producing the greenhouse gas \(\text{CO}_2\). Therefore, the development of new chemical methods to incorporate nitrogen atoms into organic molecules without the need for reduction of \(\text{N}_2\) to ammonia would effect both a more efficient and environmentally benign production of nitrogen-containing organic molecules. For these reasons, this area of research is under active exploration.14–19

In order to bypass traditional schemes for nitrogen fixation, it is important that the product of \(\text{N}_2\) transfer be a useful synthetic precursor for organic reactions. One class of compounds that meets this criteria is that of the diazoalkanes, which are important precursors to nitrogen-containing organic molecules. These molecules react by 1,3-dipolar addition reactions to incorporate both nitrogen atoms into organic functional groups.20–22 Thereby, diazoalkanes are used as synthons for pyrazolines,23–27 compounds used in asymmetric ligand25 and drug design.28–29

Diazooalkane complexes may be made from coordinated \(\text{N}_2\), such as one \(\text{N}_2\) ligand in \((\text{N}_2)_2\text{M(diphos)}_2 (\text{M} = \text{Mo}, \text{W}; \text{diphos} = \text{chelating diphosphine})\),1,17,20,30–32 or by treatment of a transition-metal complexes with an organic diazoalkane to form a coordination complex.33–40 For instance, photolysis of gem-dibromides (i.e., \(\text{Me}_2\text{CBr}_2\)) in the presence of \((\text{N}_2)_2\text{W(diphos)}_2\) has been shown to afford \([\text{Me}_2\text{CNNW(diphos)}_2]\text{Br}\).31 An alternative route to diazoalkane complexes of this type has been accomplished by the sequential treatment of \((\text{N}_2)_2\text{M(diphos)}_2\) with a strong acid followed by an aldehyde. In those diazoalkane complexes that are prepared by addition of an organic diazoalkane to a transition-metal complex, either hydrazones or diazenes may serve as precursors to the diazoalkane ligand.41–46 In many cases, compounds arrived at by either synthetic route are stable with regard to \(\text{N}_2\) loss that would lead to formation of a carbene complex.1,17,20,35–46 However, diazoalkanes also remain synthons of choice to generate metal carbenes.33–37,44–52

Given that diazoalkane complexes may be made by addition of a diazoalkane to a coordinatively unsaturated transition metal complex, one might expect that \(\text{Ph}_2\text{CN}_2\) would add to \(\text{Mo(N}[\text{t-Bu}\text{Ar}])_3\) to form a diazoalkane complex, in analogy to the odd-electron compound \(\text{PhCNM}o(\text{N}[\text{t-Bu}\text{Ar}])_3 (\text{Ar} = 3,5-\text{C}_6\text{H}_2\text{Me}_2).53–55\)
However, we found that both Ph₂CN₂ and N₂O engage in facile reactions with Mo(N[R-Bu]Ar)₃ that form NMо(N[R-Bu]Ar)₃.⁵⁶,⁵⁷ As a result, odd-electron diazoalkane complexes cannot be prepared by simple coordination of a free diazoalkane to Mo(N[R-Bu]Ar)₃. This finding motivated us to develop alternative routes for the synthesis of the complexed diazoalkane functional group in order to prepare complexes of the formula 4-RC₆H₄C(H)NNMo(N[R-Bu]Ar)₃, R (R = NMe₂, Me, H, Br, CN).

The N₂-derived diazenido compound, Me₃SiNNMo(N[R-Bu]Ar)₃ is formed in high yields by treatment of Mo(N[R-Bu]Ar)₃ with a mixture of 1% Na/Hg and Me₃SiCl under an atmosphere of N₂.⁵⁸ We thought that the terminal nitrogen of the trimethylsilyl diazenido would be sufficiently nucleophilic to participate in reactions that form N–C bonds, and therefore, protocols were developed to convert this diazenido fragment into a coordinated diazoalkane.

Results and Discussion

Synthesis and Structure of Diazoalkane Complexes. Inspired by organic condensation reactions between aldehydes and amines,⁵⁹ we explored treatment of Me₃SiNNMo(N[R-Bu]Ar)₃ with aldehydes. A related strategy has been successful in the case of [H₂NNM(diphos)₂F][BF₄] (M = Mo, W), which undergoes reactions with ketones to lose 1 equiv of H₂O forming [R₂CNNM(diphos)₂F][BF₄].¹⁷,³⁰,³² Because Brønsted acids are incompatible with our ancillary ligands, strong Lewis acids were used in place of protic conditions. Accordingly, Me₃SiNNMo(N[R-Bu]Ar)₃ was treated with 1 equiv of AlCl₃ and PhCHO in Et₂O, upon which the product diazoalkane complex [PhC(H)NNMo(N[R-Bu]Ar)₃][AlCl₄], [1-H][AlCl₄], precipitates from the reaction mixture as a red powder (Scheme 1). However, half of the starting diazenido complex was recovered from the reaction mixture when this procedure was followed. When 2 equiv of AlCl₃ were used, [1-H][AlCl₄] was isolated from the reaction mixture in 90% yield by filtration. We presume that the soluble byproduct of this synthesis is [μ-(Me₃SiO)AlCl₂].⁶₀,⁶¹ The salt, [1-H][AlCl₄], is stable with regard to N₂ loss. In fact, [1-H][AlCl₄] may be heated to 60 °C in CDCl₃ for several days without noticeable decomposition (as assayed by ¹H NMR), and solids may be repeatedly melted at 192 °C.

Employing N₂ as the synthon for the diazoalkane ligand of [1-H][AlCl₄] made it convenient to isotopically label the diazoalkane fragment via use of ¹⁵N₂. The ¹⁵N NMR spectrum of ¹⁵N₂-[1-H][AlCl₄] depicts two doublets of doublets centered at 384 and 417 ppm. These values agree well with the chemical shifts of 397 and 422 ppm obtained from DFT calculations (Table S1). The deshielding of these nitrogen nuclei has been attributed to low-energy nN → π* circulations.⁶² The observed ¹JNN = 17 Hz coupling is consistent with literature data for metal diazenido compounds in which the metal-N-N angle is approximately linear.⁶² We assign the more downfield signal as the nitrogen bound to Mo based upon both proton coupling and comparison to related compounds such as Me₃SiNNMo(N[R-Bu]Ar)₃. The proton-coupled ¹⁵N NMR spectrum revealed N–H couplings of ³JNNH = 6 Hz and ²JHH = 3 Hz consistent with the expectation that the magnitude of the ²JHH coupling through an sp² hybridized atom will be smaller than that of the ³JNNH coupling for a planar diazoalkane ligand.⁶³

The synthesis of [1-H][AlCl₄] requires both that an oxygen atom be removed from the aldehyde, breaking the C=O double bond, and that a C=N double bond be formed during the reaction. It is also a redox reaction in which molybdenum goes from the +4 to +6 formal oxidation state. We found it interesting that such a transformation gave a diazoalkane-containing product in high yields. Moreover, the thermal stability of [1-H][AlCl₄] contrasts

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**Scheme 1**

<table>
<thead>
<tr>
<th>R</th>
<th>σ</th>
<th>Yield (%)</th>
</tr>
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<tr>
<td>NMe₂</td>
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<td>51</td>
</tr>
<tr>
<td>Me</td>
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<td>0.66</td>
<td>72</td>
</tr>
</tbody>
</table>

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*Synthesis of diazoalkane complexes. Yields and Hammett σ parameters for the aryl substituent are given in the Scheme.*

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**Figure 1.** An ORTEP drawing of [1-H][AlCl₄] with 50% probability ellipsoids. Hydrogen atoms having calculated atomic positions are omitted for clarity. Selected bond lengths and angles are given in the text.
with previous use of this diazoalkane, PhC(H)N2, to study carbene transfer reactions which occur via N2 loss. Therefore, we sought to expand the synthetic methodology in a manner suitable for systematic comparison of the properties of such diazoalkane complexes. For this purpose, we chose to vary the substituent in the 4-position of the diazoalkane aryl ring. This strategy allows us to draw upon linear free energy relationships established within the field of physical organic chemistry as a guide for comparison. We found that with only minor modifications in the protocol, the synthesis of [1-H][AlCl4] could be successfully applied for the synthesis of four other diazoalkane complexes [4-RC6H4C(H)NNMo(N[t-Bu]Ar)3][AlCl4], [1-R][AlCl4] (R = NMe2, 51%; Me, 90%; Br, 97%; CN, 72% isolated yield; Scheme 1) by combination of the appropriate aldehyde, AlCl3, and Me3SiNNMo(N[t-Bu]Ar)3.

Figure 2. An ORTEP drawing of [1-NMe2][AlCl4] with 50% probability ellipsoids. Hydrogen atoms having calculated atomic positions are omitted for clarity. Selected bond lengths and angles are given in the text.

Crystals of [1-H][AlCl4] suitable for X-ray diffraction were grown from THF/n-pentane at –35 °C. The X-ray crystal structure of [1-H][AlCl4] shows that the three anilide ligands are oriented in the same direction, imparting a pseudo-C3 symmetry to the lower hemisphere of the molecule that is broken by the Cs symmetric diazoalkane ligand (Figure 1). The Mo–N4 distance of 1.7326(18) Å suggests π bonding between Mo and the diazoalkane N. The N4–N5 distance of 1.336(2) Å and N5–C4 distance of 1.290(3) Å are consistent with the description that the diazoalkane ligand is reduced by backbonding from Mo. Accordingly, the N–N distance is substantially longer than what one would expect for a free diazoalkane (cf. 1.13(4) Å for H2CN2). A slightly different set of internuclear distances were found in the single-crystal X-ray structure of [1-NMe2][AlCl4]: Mo–N4, 1.730(2) Å; N4–N5, 1.317(3) Å; and N5–C4, 1.312(4) Å (Figure 2). The NMe2 group is coplanar with the aryl ring, maximizing the electron donation of this substituent into the π network of the diazoalkane. Presumably, this electronic perturbation is responsible for the small alteration in internuclear distances in [1-NMe2][AlCl4] compared to [1-H][AlCl4]. Both sets of metrics compare well with those obtained for other terminally bound, singly-bent diazoalkanes coordinated to Mo and W. Diazoalkane ligands are described as singly-bent when the C–N–N angle is approximately 120° and the metal–N–N angle is close to 180°, as is the case for the compounds reported here. The N4–N5–C4 and Mo1–N4–N5 angles are 117.86(19)° and 174.21(16)° in [1-H][AlCl4] and 116.9(3)° and 176.0(2)° in [1-NMe2][AlCl4].

Figure 3. IR (above) and Raman (below) spectra of [1-H][AlCl4] (solid blue line) and 15N2-[1-H][AlCl4] (dashed green line) in CDCl3 at 20 °C.

Electronic and Vibrational Spectroscopy. Diazoalkane complexes typically have characteristic stretching modes that are suitable for study by vibrational spectroscopy. To gain information about the vibrational structure of the coordinated diazoalkane ligand, infrared and Raman spectroscopy data were collected for complexes [1-R][AlCl4] (R = NMe2, Me, H, Br, CN). Due to the low symmetry of the five cations (approximately Cs), most vibrational modes of the diazoalkane ligand give rise to bands in both the Raman and IR spectrum (Figures S15-S20). The Raman
spectrum of [1-H][AlCl4] contains five intense bands that are attributed to vibrational modes of the PhC(H)N2 ligand located at 1528, 1446, 1366, 1193, and 1173 cm–1. The band located at 1528 cm–1 in both the IR and Raman spectra is in the spectral region for which the predominantly C=N stretching mode of the η1-diazoalkane ligand is expected. This assignment was supported by a 15N2-labeling experiment in which the corresponding band for 15N2-[1-H][AlCl4] was found at 1520 cm–1 (Figure 3). The 8 cm–1 shift of the C=N stretching mode is smaller than would be predicted by a harmonic oscillator model (24 cm–1), indicating that this vibrational mode couples to others. Coupling of the C–N and N–N vibrational modes is observed for both coordinated and free diazoalkanes. Additionally, for aryl diazenido complexes it has been demonstrated that the aryl C–H modes couple into the N–N oscillator.

Table 1. Vibrational data for the diazoalkane ligand in cm–1.

<table>
<thead>
<tr>
<th>Complex</th>
<th>ν1</th>
<th>ν2</th>
<th>ν3</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1498</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[1-Me][AlCl4]</td>
<td>1526</td>
<td>1367</td>
<td>1197</td>
</tr>
<tr>
<td>[1-H][AlCl4]</td>
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<td>1520</td>
<td>1346</td>
<td>1164</td>
</tr>
<tr>
<td>[1-Br][AlCl4]</td>
<td>1525</td>
<td>1367</td>
<td>1192</td>
</tr>
<tr>
<td>[1-CN][AlCl4]</td>
<td>1524</td>
<td>1361</td>
<td>1193</td>
</tr>
</tbody>
</table>

* Sample fluorescence prevented the acquisition of Raman data.

A band that is only intense in the Raman spectrum of [1-H][AlCl4] was located at 1366 cm–1. The Raman activity of this band suggested to us that it is largely composed of N–N stretching character which is expected to give rise to a smaller change in dipole moment than the C=N stretching motion. When 15N2 is incorporated into the diazoalkane ligand this band shifts to 1346 cm–1. The 20 cm–1 shift that occurs upon isotopic substitution is twice as large as the shift found for the predominantly C=N stretching mode, but is less than half the predicted value of 46 cm–1 for a harmonic N–N oscillator. Calculations carried out on [1-m]+ found a band at 1363 cm–1 that is described by a mixture of both the C–N and N–N stretching modes combined with a wagging of the hydrogen atom attached to the diazoalkane carbon. For this reason, the band at 1366 cm–1 is attributed to a highly coupled vibrational mode that contains both C=N and N–N stretching character.

In the Raman spectrum of [1-H][AlCl4], the two bands at 1193 and 1173 cm–1 for collapse into a single band located at 1164 cm–1 upon 15N substitution. We suggest that this is because the two signals overlap in the spectrum of the labeled compound. The band located at 1193 cm–1 is both sharp and intense in the Raman spectrum, consistent with an N–N stretching mode. Moreover, this band shifts by 29 cm–1 upon isotopic substitution compared with the 40 cm–1 shift predicted by a harmonic oscillator model, further supporting the notion that the origin this band is a stretching mode of the N2 unit. DFT calculations support the description of this vibration as a predominantly N–N stretching mode. For [1-m]+ a band of this type was found at 1148 cm–1 which is in close agreement with experimental data given the simplicity of the model system.

Given the wealth of data available for the IR-active, C=N stretching mode of coordinated diazoalkanes, information for the N–N stretching mode remains sparse in comparison to information concerning the same stretch for free diazoalkanes. The N=N stretching mode of PhC(H)N2 is 2062 cm–1 which is between the values for H2CN2, 2097 cm–1, and Ph2CN2, 2042 cm–1. The observed N–N stretch in [1-H][AlCl4] is consistent with a weakening of this bond in comparison to the free diazoalkane.
All of the diazoalkane complexes studied have similar vibrational structure (Table 1). In view of the isotopic substitution results obtained for [1-H][AlCl4], the predominantly C=N and N–N stretching modes were located in the IR and Raman spectra for each of [1-R][AlCl4] (R = NMe2, Me, Br, CN; Table 1). There is no obvious Hammett trend present in the vibrational data, which may reflect the coupling of the multiple vibrational modes. Vibrational modes that have weak intensity in the infrared spectrum were not located for [1-NMe2][AlCl4] because sample fluorescence interfered with the acquisition of Raman data.

The compounds [1-R][AlCl4] (R = Me, H, Br, CN) are all dark red in color, but solutions of [1-NMe2][AlCl4] are a vibrant fuchsia color. This observation prompted us to compare the UV-Vis spectral features of these five complexes. The UV-Vis spectra for [1-R][AlCl4] (R = Me, H, Br, CN) all show an absorption near 450 nm and contain no intense absorption bands at lower energies (Figure 4). However, [1-NMe2][AlCl4] was found to possess an intense absorption with maximum at 538 nm (Figure 5). The absorption band is assigned as a ligand-to-metal charge transfer (LMCT) band (LMCT). This assignment is in accord with both the formally intense absorption with maximum at 538 nm (Figure 5). The absorption may be fit to the sum of three Gaussian distributions with spacing of approximately 1180 cm⁻¹. This value likely corresponds to the vibrational mode of the Mo–N bond in the excited state.80,81

Irradiation of CHCl₃ solutions of [1-NMe2][AlCl4] with visible light gave rise to an emission with a Stokes shift of 15600 cm⁻¹ (λₑₓ = 541 nm, λᵣₑᵢ = 700 nm). A number of workers have recently shown that the LMCT excited state may give rise to fluorescence in d⁰ complexes that contain metal-ligand multiple bonds.82-84 Our case is distinct in that both absorption and emission of the LMCT excited state occur in the visible region, therefore UV irradiation is not required to observe emission from [1-NMe2][AlCl4]. Indeed, the NMe₂ substituent is found in many efficient organic fluorophores that emit from a CT state.85 This is desirable for applications such as photooxidants that may be developed for use with a visible light source.86

Reductive Dimerization of Diazoalkane Ligands via C–C Bond Formation. Reduction of the diazoalkane salt, [1-H][AlCl4] by 1e was anticipated to generate the odd-electron complex PhC(H)NNMo(N[t-Bu]Ar)₂, 1-H. However, this species was found to be unstable with regard to spontaneous dimerization at carbon to form the dimeric, diazo complex, μ-(PhC(H)NN)₂[Mo(N[t-Bu]Ar)₂]₂, [1-H]₂ (Scheme 2). Addition of solid Cp₂Co to [1-H][AlCl4] dissolved in CHCl₃ leads to the rapid and quantitative formation of [1-H]₂; with precipitation of [Cp₂Co][AlCl₄]. Filtration of the reaction mixture removes [Cp₂Co][AlCl₄], leaving only [1-H]₂ in the filtrate which is then collected in 97% yield. This was found to be the most convenient protocol for separating the products despite the observation that Cp₂Co is unstable in CHCl₃ for prolonged periods of time; the formation of [1-H]₂ is essentially complete upon mixing. Dimeric [1-H]₂ has low solubility in solvents such as C₆H₆, Et₂O, or n-alkanes but forms solutions readily in THF, MeCN, or halocarbons. Upon reduction of [1-H][AlCl₄], the characteristic downfield shift at 8.92 ppm in the ¹H NMR spectrum for the hydrogen atom bound to the diazoalkane carbon moves to 6.17 ppm. This indicates that the unsaturation at that carbon is removed upon reduction, consistent with dimer formation. Examining the ¹H NMR spectrum, it intrigued us that only one set of resonances predominated; one would expect an unselective radical-radical coupling reaction to form a 1:1 mixture of rac and meso isomers with each isomer having a unique set of chemical shifts. To rationalize this observation, we concluded that the reductive dimerization of [1-H][AlCl₄] is selective for the formation of either the meso or rac product mixtures.

To identify the preferred stereochemistry of C–C bond formation, crystals suitable for X-ray diffraction were grown of [1-H]₂. The molecule crystallizes on a C₂ axis and is therefore assigned as the rac isomer (Figure 6). Anomalous scattering data confirmed that the crystallographic model with (R, R) stereochemistry corresponds to the enantiomer contained within the crystal.86-88 The preferential formation of the rac isomer is in qualitative agreement with a radical-radical coupling in which a transition state leading to the rac product is favored. This is the only transition state that accommodates C–C bond formation with C₄₁–C₄–C₄¹‘–C₄₁’ and N₅–C₄–C₄¹‘–N₅¹’ angles that approximate 180°, minimizing unfavorable steric interactions between the two reaction partners. Likewise, reduction of both [1-Br][AlCl₄] or [1-CN][AlCl₄] with Cp₂Co in d₅-THF leads to the formation of only one set of signals in the ¹H NMR spectrum which we presume to be the rac isomer as for [1-H]₂. However, reduction of [1-Me][AlCl₄] or [1-NMe₂][AlCl₄] gives mixture of the meso and rac isomers as judged by ¹H NMR spectroscopy.
Highly selective formation of the rac product for the dimerization of 1-R (R = H, Br, CN) is consistent with a number of mechanisms. If C–C bond formation were reversible, an equilibrium process involving [1-R]_2 and 2 equiv of 1-R could drive the dimer to the more stable stereoisomer. However, based on the large difference in $E_P$ for [1-R]^{0+} and [1-R]^{0+} of approximately 900 mV (vide infra), this explanation appears unlikely. A related scenario is that formation of [1-R]^{2+} by combination of 1-R and [1-R]^{+} is rapid and reversible with respect to the second electron transfer. In such a mechanism, the stereochemical outcome of C–C bond formation might be explained by the relative stability of the two stereoisomers of [1-R]_2. An equally plausible possibility is that the two neutral radicals, 1-R, couple in such a way as to minimize steric repulsion which leads to a preferential formation of rac isomers. Electrochemical studies of the pinacol coupling involving 4-substituted acetophenones show a modest preference for the formation of the rac isomers in basic media. Although these results were attributed mainly to hydrogen bonding, similar electrodimerization of benzaldehyde did not discriminate between formation of rac or meso isomers.

With respect to [1-H][AlCl_4], the C–C bond formation forces the cumulene system to reverse the long and short internuclear distances in [1-H]_2. This structural change occurs in conjunction with the conversion of molybdenum from the +6 to the +4 oxidation state in the reaction during the reductive coupling. The Mo–N distance in [1-H]_2, 1.769(2) Å, is longer than that of [1-H][AlCl_4]. Correspondingly, the N–N bond is contracted, 1.243(4) Å, and the C–N bond lengthened, 1.478(4) Å, compared to the cationic complex. The newly formed C–C linkage has an internuclear distance of 1.553(6) Å which is typical for such a C–C single bond. In the X-ray crystal structure of [1-H]_2, the N–N–C bond angle of 119.4(3)° is consistent with $sp^2$ hybridization at nitrogen, as the Mo–N–N angle of 167.2(3)° deviates from linear. This angle is expected to be less than 180° in metal diazenido complexes. Hoffmann has described the bonding in this functional group (generically, M–N–N–H) and found a dependence of the metal–N–N and N–N–H angles. These calculations concluded that when the latter angle is 120°, the M–N–N angle is predicted to be 174°.

Electrochemical dimerization of coordinated diazoalkanes has been reported for two systems that contain W, eq. 1. In only one case was the product soluble enough for spectroscopic and structural characterization, eq. 1. In this case, reduction of [Me_2CNNWCl(CO)(PMe_2Ph)_3]^+, is accompanied by loss of a phosphine ligand, therefore the reaction was carried out in the presence of CO to cleanly generate the C–C bonded product. Moreover, there is no possibility of forming diastereomeric products because the two faces of the diazoalkane ligand are identical. Reductive dimerization of diazoalkane complexes has been reported for two systems that contain W, eq. 1. In only one case was the product soluble enough for spectroscopic and structural characterization, eq. 1.
Figure 7. Potentials ($E_{1/2}$ vs. SCE) at which reduction of [1-R][AlCl₄] occurs are plotted against the Hammett parameter for R. The fit shown has a $\rho = -283$ mV ($R^2 = 0.998$).

Figure 8. Stack plot of cyclic voltammograms of 1 mM [1-H][AlCl₄] in 0.2 M [N(n-Bu)₄][PF₆]/THF solution at 20 °C. Potentials shown are relative to SCE.

In cyclic voltammetry, the ratio of cathodic-to-anodic peak currents reflects the concentration in solution for the generation and decay of electroactive species. Therefore, monitoring this ratio as a function of scan rate is a sensitive marker of how much electroactive species is in solution. The height of the cathodic wave that produces 1-H varies linearly with the square-root of scan rate – as expected for a 1e reduction (Figure 8). However, the anodic current corresponding to oxidation of 1-H grows in more quickly than predicted by the scan rate variation. This behavior shows that in 1 mM solution 1-H is long-lived enough to be oxidized prior to dimerization at sufficiently fast scan rates. The low anodic peak current for slow scan rates indicates that depletion of 1-H in solution, via dimerization, is fast relative to the timescale of the electrochemical experiment. Consistent with a bimolecular dimerization reaction, cyclic voltammograms of [1-R][AlCl₄] were concentration dependent. Increasing the concentration led to less reversible waves while decreasing the concentration gave more reversible behavior.

Solutions of [1-R][AlCl₄] (R = Br, H, Me) at concentrations of 1 mM have nearly the same cathodic peak current, indicating that the diffusion behavior of the three compounds is similar. A variable scan rate experiment showed that the ratio of cathodic-to-anodic peak current is not equal for the series (Figure 9). Upon comparing the cyclic voltammograms of [1-H][AlCl₄] and [1-Me][AlCl₄] under the same conditions (1 mM solutions at a scan rate of 100 mV/s), the wave corresponding to [1-Me]$^+$ appears more reversible than that for [1-H]$^+$ because the ratio of cathodic-to-anodic peak currents is closer to unity. Assuming that the odd-electron diazoalkane complexes, 1-R, have similar diffusion behavior, we would conclude that this ratio should be the same for all three substituents. This result demonstrates that the dimerization process is not diffusion limited. The dimerization reaction that forms 1-H₂ as a product by reduction of 1-H$^+$ has two limiting mechanisms. Two equiv of the neutral, odd-electron compound 1-H may dimerize forming a C=C bond. Alternatively, 1-H may react with 1-H$^+$ to form an intermediate complex 1-H$^+$. This mechanism may be discriminated by monitoring the changes in the cathodic peak potential as a function of scan rate. In the radical-radical coupling case a slope of –19 is expected for such a plot while a slope of –29 is expected for a radical-cation coupling. Our attempts to make this distinction were not reliable, presumably due to the high resistance of anhydrous media. Except for in the immediate vicinity of the electrode surface, the concentration of 1-H$^+$ is always expected to be higher than the concentration of 1-H, raising the question as to whether chemical reduction of these cations would necessarily follow the same mechanism that predominates under electrochemical conditions. We favor the notion that this dimerization proceeds by radical-radical coupling, by analogy with the well-studied radical-radical dimerization of the persistent radical, PhCNMoO[N(n-Bu)Ar]₃, but do not rule out a mechanism in which 1-R₂ is formed by a radical-cation dimerization followed by fast electron transfer.

Figure 9. Plots of cathodic/anodic peak currents vs. log of the scan rate (mV/s) show that the dimerization is not diffusion limited. Fits to the data in the form $i_c / i_a = a + b \log(\nu)$, where $\nu$ is the scan rate, are shown as follows: [1-Br][AlCl₄] (dashed line), $a = 7.56(26)$, $b = -1.69(10)$; [1-H][AlCl₄] (solid line), $a = 5.95(10)$, $b = -1.46(4)$; [1-Me][AlCl₄] (dotted line), $a = 3.67(6)$, $b = -0.76(3)$.

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Savéant has described the electrodimerization of aromatic aldehydes in great detail. The mechanism of dimerization has been shown to be a radical-radical coupling of two pinacolate radical anions generated at the electrode. The reaction proceeds at rates much slower than the diffusion limit, and specific solvation by water has a large effect on the measured rate of this reaction.\(^{105}\) The activation barrier to this reaction has been attributed to entropy requirements in the C–C bond forming step.\(^{96,102,103}\) A reduced spin density on the carbon atom brought about by delocalization of the unpaired electron onto the aromatic group both stabilizes the intermediate and raises the entropic requirement for C–C bond formation. Therefore, increasing the amount of radical character at the carbon participating in C–C bond formation generally increases the rate of dimerization.

The transient radical intermediate, 1-H, is unstable with respect to dimerization to form a C–C bond in the product [1-H]_2. Dimerization at carbon suggested to us that the unpaired electron resides partially on the diazoalkane carbon atom. To investigate the electronic structure of the transient radical intermediate, DFT calculations were performed on the model complex PhC(H)NMMo(NH_2)_3, 1-m. For a starting geometry the half-molecule that occupies the asymmetric unit in the crystal structure of [1-H]_2 was used and the anilide ligands were replaced with NH_2 donors. During the geometry optimization, the initially pyramidal diazoalkane carbon atom became planar as expected for an sp\(^2\) hybridization at carbon. In 1-m, the SOMO is destabilized by approximately 1.5 eV with respect to the SOMO–1 (Figure 10).

Approximately 15% of the SOMO resides on the diazoalkane carbon atom and 43% on Mo (Figure 11). Experimental evidence for the delocalization of unpaired spin density onto diazoalkane ligands has been provided by study of the reduction of [4-RC_6H_4C(H)NNMo(dppe)_2F][BF_4] in an EPR probe.\(^{95}\) An EPR resonance centered at \(g = 2.01\) was split into a quintet by hyperfine interaction of 2.6 mT from \(^{14}\)N nuclei. Delocalization of the SOMO of 1-m onto the aryl ring suggests that a para substituent could influence the stability of the SOMO and the relative spin densities for atoms along the diazoalkane fragment. This provides a model for the variation in reduction potential and lifetime of 1-R with variation of the substituent.

**Figure 10.** Relative energies of the calculated frontier orbitals for 1-m.

**Figure 11.** Two views of the SOMO for 1-m shown at an isosurface value of 0.05.

Electrochemical oxidation of the dimers [1-R]_2 gives a broad anodic wave in the cyclic voltammogram with a peak current approximately twice the magnitude of the wave corresponding to the cathodic [1-R]^−/0 couple (Figure 12). Therefore, the oxidation of [1-R]_2 was assigned as a 2e wave that occurs as a result of two consecutive 1e oxidation events.\(^{96}\) Behavior of this kind is expected for an electrochemical oxidation in which the second electron is removed more easily than the first.\(^{106}\) Consistent with this reasoning is the difference between the oxidation potential of [1-R]_2 and that of the neutral 1-R, as the potential of the latter lies approximately 1 V negative of the former. A frequently invoked mechanism for reactions of this type is the ECE reaction, wherein oxidation of the dimer, [1-H]_2, leads to formation of both [1-H]^+ and 1-H at a potential at which the neutral compound undergoes rapid oxidation to form a second equiv of [1-H]^+.\(^{99,106}\)

Interested in the electronic effects of the para substituent, we investigated oxidative cleavage for [1-R]_2. One would expect that the electronic influence of the substituent to be less dramatic in the case of dimer oxidative cleavage than was observed for reductive coupling of [1-R]_2 because of the absence of a conjugated \(\pi\) network enabling communication between the aromatic ring and molybdenum. The potential \((E_p)^{+}\) for oxidative cleavage of [1-R]_2 was measured by differential scanning voltammetry and fit to a Hammett plot with a \(\rho = –160\) mV for four points (R = Me, H, Br,
The potential for oxidation of [1-NMe$_2$]$_2$ was nearly that of [1-CN]$_2$. In order to test for a V-shaped Hammett behavior, the compound [1-OMe][AlCl$_4$] was made and briefly investigated. Oxidation of the dimeric species [1-OMe]$_2$ occurred at $E_0 = 417$ mV. This value for R = OMe fit well to a line when plotted with the other four points (R = Me, H, Br, CN). It is unclear why the potential for [1-NMe$_2$]$_2$ does not fit to the Hammett trend, however it is noteworthy that this compound has other nearby oxidation events at more positive potentials that may correspond to oxidation of the dimethylamine functional group (Figure S23).

![Figure 13. Potentials ($E_0$ vs. SCE) at which oxidative cleavage of [1-R]$_2$ occur are plotted against the Hammett parameters for R. The fit to four data points has a $\rho = -160$ mV ($R^2 = 0.994$). Data for [1-NMe$_2$]$_2$ was not fit to the line.](image)

After the transient radical I-H dimerizes to form a C=C bond, 2e are formally stored in a C=C bond. We sought to engage these electrons in a reaction that would return the starting material, [1-H]. Accordingly, oxidative cleavage [1-H]$_2$ effected by either the addition of AgOTf or [Cp$_2$Fe][PF$_6$] to [1-H]$_2$ give [1-H][OTf] (68%) and [1-H][PF$_6$] (78%) as products. This demonstrates a two-step route for contion exchange that converts the initial dimeric species [1-H][AlCl$_4$] into [1-H][A] ([A = PF$_6^-$, OTf$^-$; eq. 4, 5]).

\[
\text{[1-H][AlCl$_4$]} + \text{[Cp$_2$Co]} \rightarrow \text{[1-H]$_2$ (97%) + [Cp$_2$Co][AlCl$_4$]} \quad (4)
\]

\[
\text{[1-H]$_2$ + [Cp$_2$Fe][PF$_6$]} \rightarrow \text{[1-H][PF$_6$] (78%) + Cp$_2$Fe} \quad (5)
\]

Related examples of redox non-innocent C=C bonds have been studied for their capacity to act as charge storage devices. Metalloccumulenes such as Cp(Et$_2$PCH$_2$CH$_2$PMe$_2$)Mn=C=CH$_2$ participate in oxidative coupling reactions to form a C=C bonded dimer that formally stores two holes.\textsuperscript{107-109} The complementary function, electron storage, is accomplished by [1-H]$_2$. Combination of hole and electron storing molecules is a goal in the development of molecular batteries.\textsuperscript{109} Only a few systems have been described that carry out both functions, such as Floriani’s [Msalophen]$^n$ (n = 2, 0, –2).\textsuperscript{110-113} Therefore, it has become desirable to find mixed-molecule systems for charge storage. These systems expand the scope of molecular charge storage technology, and make it possible to tune the electrochemical difference between the hole and charge storage compartments of the molecular battery.

**Conclusions**

A new synthetic route to diazoalkane ligands from N$_2$ has been described. Characterization of the complexes, [1-R][AlCl$_4$], included Raman spectroscopy to elucidate the N-N stretching frequency of the coordinated diazoalkane, for which literature data are scarce. Reductive dimerization of [1-R] coupled the diazoalkane ligands at carbon to form a bridging ligand. One such dimer, [1-H]$_2$ was crystallographically characterized. Electrochemical studies showed that reduction of [1-R][AlCl$_4$] forms a transient odd-electron complex, 1-R. The structure of this odd-electron intermediate was probed using quantum chemical calculations. Future work might be aimed at liberation of the coordinated diazoalkane from the metal scaffold and the application of C=C coupled diazoalkane ligands in charge storage devices.

**Experimental Procedures**

**General Remarks.** All manipulations were carried out under an atmosphere of purified nitrogen in a Vacuum Atmospheres Model MO-40M glovebox equipped with the QP-30 accessory, or by standard Schlenk techniques.\textsuperscript{114,115} Inside the MO-40M glovebox the ambient temperature ranged from 18–22 °C. All glassware was oven-dried at a temperature above 150 °C for at least 12 hours and allowed to cool under dynamic vacuum prior to use. Celite, alumina, and 4 Å sieves were activated by heating to a temperature greater than 180 °C under a dynamic vacuum for 2 d (Celite) or 5 d (alumina and 4 Å sieves). Et$_2$O, n-hexane, n-pentane, and toluene were bubble degassed with nitrogen and forced, under positive pressure, through a column of activated alumina followed by a column of activated silica gel. MeCN was taken from an Aldrich Sure-Seal bottle, filtered through a column of activated alumina (4 × 3 cm), and degassed under a dynamic vacuum. O(SiMe$_3$)$_2$ was distilled from dark purple solutions of sodium benzophenone ketyl. MeCN was taken from an Aldrich Sure-Seal bottle, filtered through a column of activated alumina (4 × 3 cm), and degassed under a dynamic vacuum. (SiMe$_3$)$_2$ was distilled from dark purple solutions of sodium benzophenone ketyl. CHCl$_3$ was refluxed over CaH$_2$ for 48 h prior to distillation. All solvents were stored over 4 Å sieves. C$_2$D$_6$ was degassed by three freeze-pumpthaw cycles and stored over 4 Å sieves for 3 d prior to use. CDCl$_3$ and CD$_2$Cl$_2$ were refluxed over CaH$_2$ for 24 h then distilled and stored over 4 Å sieves. $^1$H and $^{13}$C NMR shifts are referenced to residual solvent resonances (for CD$_2$Cl$_2$ δ = 49.3 ppm and 54.00 ppm). All $^{15}$N NMR spectra are externally referenced to neat H$_2$C=C$^{15}$N (δ = 245 ppm) in comparison to liquid NH$_3$ ($\delta = 0$ ppm).\textsuperscript{117} UV-Vis spectra were obtained on HP8453 spectrophotometers in 1 cm quartz cells manufactured by Starna. Infrared spectra were obtained on a Nicolet Magna 860 FT-IR in transmission mode. Positive ion ESI-MS were obtained using a Bruker Daltonics APEXIV, 4.7 T Fourier Transform Ion Cyclotron Resonance Mass Spectrometer. Combustion analysis was performed by Midwest Microlab, LLC (Indianapolis, IN).

**Raman Spectroscopy.** An Invictus solid state laser at 785 nm manufactured by Kaiser Optics was routed through fiber optic cables to a Hololab series 5000 Raman Microscope. The sample was placed in an NMR tube under the objective (10×) of the microscope. The Stokes shifted Raman scattering was observed via
180° reflectance through the objective of the Raman microscope at a laser power of 30 mW. Reported Raman spectra are an average of 8 spectra, each collected for 2 s using an automated routine in Hololab software to automatically correct for the dark current. The averaged spectrum has the solvent signals subtracted and was subsequently baseline corrected using GRAMS software. Raman shifts are reported in reference to an external standard of neat cyclohexane (801.3 cm⁻¹).

**Electrochemical Measurements.** A Bioanalytic Systems CW-50 potentiostat, 1 mm diameter Pt disk working electrode, Pt wire counter electrode, and Ag wire pseudo-reference electrode were used for all measurements. All electrochemical measurements were made in 0.2 M [N(n-Bu)4][PF6] THF solutions. To measure potentials, Cp*Fe was used as an internal standard. The Cp*Fe²⁺ is referenced to 171 mV vs SCE. Reversible redox potentials of [1-R][AlCl4] were measured by cyclic voltammetry of 50 μM solution at several scan rates. Oxidative peak potentials of in situ formed [1-R]: were measured by differential pulse voltammetry (dpv) acquired on 5 mM solutions of [1-R][AlCl4]. DPV data were acquired with a single scan from ca. –1 V to 1 V taken at 20 mV/s using the following data collection parameters: pulse height, 50 mV; pulse length, 50 ms; sampling intervals, 20 ms; and cycle repeat time, 200 ms. For R = OMe, all potentials were measured at arbitrary concentrations. Isolated samples of [1-H][AlCl4] and [1-H2] show the same features in the cyclic voltammogram.

**Single Crystal X-ray Diffraction.** X-ray diffraction data were collected using a Siemens Platform three-circle diffractometer coupled to a Bruker-AXS Smart Apex CCD detector with graphite-monochromated Mo Kα radiation (λ = 0.71073 Å), performing φ- and ω-scans. The molecular structure was obtained by direct methods using SHELXS and refined against F². The 20 °C): δ = 8.56 (1 H, s, CHN2), 7.84 (2 H, br, Me2NC3H2), 6.98 (3 H, s, para-Ar), 6.81 (2 H, d, H2C(H)N2), 5.78 (6 H, s, ipso-Ar), 3.24 (6 H, s, NCH3), 2.18 (18 H, s, Ar–CH3), 1.26 (27 H, s, C(CH3)3) ppm. 13C NMR (125 MHz, CDCl3, 20 °C): δ = 163.8 (N=C), 156.1 (Me2NC), 145.5 (ipso-Ar), 138.6, 130.7, 129.3, 116.6, 112.9, 68.0 (CMe2), 40.7 (NMe2), 21.6 (ArMe ppm; one carbon was obscured. 27Al NMR (125 MHz, CDCl3, 20 °C): δ = 103.4 ppm. ESI-MS: m/z = 787.4348 (787.4339, M⁺). Anal. Cald for C44H52AlCl4N5O: C, 56.61; H, 6.84; N, 8.85; Cl, 14.85. Found: C, 56.42; H, 6.84; N, 8.85; Cl, 14.85. UV-Vis: λ (ε) = 538 (45400), 509 (35300, shoulder) nm (M⁺ cm⁻¹). MP = 125-135 °C (dec).

**Preparation of [4-RC4H4C(H)NNMo(N[R-Bu]Ar)][AlCl4], [1-R][AlCl4].** The procedure used for [1-H][AlCl4] is given. Minor modifications of this procedure were used to in the synthesis of all other diazoalkane salts, [1-R][AlCl4], discussed herein. See Supporting Information for complete preparative details.

**Characterization Data for [4-MeNC4H4C(H)NNMo(N[R-Bu]Ar)][AlCl4], [1-NMe2][AlCl4].** H NMR (600 MHz, CDCl3, 20 °C): δ = 8.78 (1 H, s, CHN2), 7.84 (2 H, br, Me2NC3H2), 7.10 (2 H, d, MeOC2H2), 7.00 (3 H, s, para-Ar), 5.74 (6 H, s, ipso-Ar), 3.94 (3H, s, OCH3), 2.18 (18 H, s, Ar–CH3), 1.26 (27 H, s, C(CH3)3) ppm. 13C NMR (125 MHz, CDCl3, 20 °C): δ = 166.4 (MeOC), 146.8 (N=C), 145.2 (ipso-Ar), 138.7, 133.4, 130.9, 128.9, 122.0, 115.8, 68.8 (CMe2), 56.2 (OMe2), 32.0 (CMe2), 21.3 (Ar-Me ppm; one carbon was obscured. 27Al NMR (125 MHz, CDCl3, 20 °C): δ = 102.8 ppm. ESI-MS: m/z = 787.4395 (787.4339, M⁺). Anal. Cald for C44H52AlCl4N5O: C, 56.61; H, 6.84; N, 8.85; Cl, 14.85. Found: C, 56.42; H, 6.84; N, 8.85; Cl, 14.85. UV-Vis: λ (ε) = 538 (45400), 509 (35300, shoulder) nm (M⁺ cm⁻¹). MP = 125-135 °C (dec).

**Characterization Data for [4-MeOC4H4C(H)NNMo(N[R-Bu]Ar)][AlCl4], [1-OMe][AlCl4].** H NMR (600 MHz, CDCl3, 20 °C): δ = 8.78 (1 H, s, CHN2), 8.03 (2 H, d, H2C(H)N2), 7.10 (2 H, d, MeOC2H2), 7.00 (3 H, s, para-Ar), 5.74 (6 H, s, ipso-Ar), 3.94 (3H, s, OCH3), 2.18 (18 H, s, Ar–CH3), 1.26 (27 H, s, C(CH3)3) ppm. 13C NMR (125 MHz, CDCl3, 20 °C): δ = 166.4 (MeOC), 146.8 (N=C), 145.2 (ipso-Ar), 138.7, 133.4, 130.9, 128.9, 122.0, 115.8, 68.8 (CMe2), 56.2 (OMe2), 32.0 (CMe2), 21.3 (Ar-Me ppm; one carbon was obscured. 27Al NMR (125 MHz, CDCl3, 20 °C): δ = 103.4 ppm. ESI-MS: m/z = 787.4384 (787.4339, M⁺). Anal. Cald for C44H52AlCl4N5O: C, 56.61; H, 6.84; N, 8.85; Cl, 14.85. Found: C, 55.86; H, 6.26; N, 7.53; Cl, 15.00%.
H, d, MeC(CH$_3$)$_2$, 7.02 (3 H, s, para-Br), 5.75 (6 H, s, ortho-Br), 2.50 (3 H, s, C$_6$H$_5$CH$_2$), 2.19 (18 H, s, Ar–CH$_3$), 1.28 (27 H, s, C(CH$_3$)$_3$) ppm. $^1$C NMR (125 MHz, CDCl$_3$, 20 °C): δ = 165.6 (N=C), 147.7, 145.3 (ipso-Br), 138.8, 131.0, 131.0, 128.9, 127.5, 69.2 (CMe$_3$), 32.1 (CMe$_3$), 22.5 (C$_2$H$_5$Me), 21.6 (Ar-Me) ppm. $^{31}$Al NMR (125 MHz, CDCl$_3$, 20 °C): δ = 103.4 ppm. ESI-MS: m/z = 758.4081 (758.4073, M$^+$. Anal. Caled for C$_{43}$H$_{60}$AlCl$_4$MoN$_5$: C, 56.62; H, 6.64; N, 7.68; Cl, 15.33%. MP = 191-193 °C.

Characterization Data for [Ph(C(H)NN)Mo(N-[t-Bu]Ar)]Cl$_4$ [1-H][AlCl$_4$]. $^1$H NMR (600 MHz, CDCl$_3$, 20 °C): δ = 8.96 (1 H, s, CF$_3$), 8.10 (2 H, d, ortho-Ph), 7.70 (1 H, t, para-Ph), 7.61 (2 H, t, meta-Ph), 7.04 (3 H, s, para-Br), 5.75 (6 H, s, ortho-Br), 2.20 (18 H, s, Ar–CH$_3$), 1.29 (27 H, s, C(CH$_3$)$_3$) ppm. $^{13}$C NMR (125 MHz, CDCl$_3$, 20 °C): δ = 165.5 (N=C), 145.3 (ipso-Br), 135.8, 131.1, 130.9, 130.9, 129.9, 128.8, 69.4 (CMe$_3$), 32.1 (CMe$_3$), 21.6 (Ar-CH$_3$) ppm. $^{31}$Al NMR (125 MHz, CDCl$_3$, 20 °C): δ = 103.4 ppm. ESI-MS: m/z = 744.3939 (744.3917, M$^+$. Anal. Caled for C$_{39}$H$_{52}$AlCl$_7$MoN$_5$: C, 56.62; H, 6.64; N, 7.68; Cl, 15.35. Found: C, 56.77; H, 6.73; N, 7.36; Cl, 15.33%. MP = 191-193 °C.

Preparation of $\mu$-[Ph(C(H)NN)Mo(N-[t-Bu]Ar)]Cl$_4$ [1-H][AlCl$_4$]. Into 1.0 mL of CDCl$_3$ was dissolved $[1$-H][AlCl$_4$] (50 mg, 0.055 mmol) to form a red solution. The solution was then heated to 50 °C for 5 h over which time no color change was observed. Analysis by $^1$H NMR spectroscopy showed only [1-H][AlCl$_4$], and no evidence of decomposition. $\mu$-[Ph(C(H)NN)Mo(N-[t-Bu]Ar)]Cl$_4$ [1-H][AlCl$_4$]. Into 1.0 mL of CDCl$_3$ was dissolved $[1$-H][AlCl$_4$] (50 mg, 0.055 mmol) to form a red solution. The solution was then heated to 50 °C for 5 h over which time no color change was observed. Analysis by $^1$H NMR spectroscopy showed only [1-H][AlCl$_4$], and no evidence of decomposition.
(4 H, br s, NCC₂H₇), 7.43 (4 H, s, H₂C₆(C(H)N₂), 6.79 (6 H, s, para-Ar), 6.32 (2 H, s, C(H)N₂), 5.78 (12 H, s, ortho-Ar), 2.10 (36 H, s, Ar–CH₂), 1.26 (54 H, s, C(CH₃)₃).

**Oxidative Cleavage of μ-(PhCH(NN)₂)Mo[N=0-Bu]Ar₂]-[H]₂ by [Cp₂Fe][PF₆].** A 5 mL THF solution of [H]-I (300 mg, 202 mmol) was frozen in a liquid nitrogen-cooled cold well. To the frozen solution was added [Cp₂Fe][PF₆] (135 mg, 408 mmol, 2 equiv) as a solid. Upon thawing of the initially yellow solution, an immediate color change to red-orange occurred. The mixture was stirred for 2 h at room temperature. The mixture was filtered through celite and the celite pad was washed with 5 mL of THF. The combined filtrate was dried under a dynamic vacuum to afford a red-orange powder. The powder was stirred in 15 mL n-pentane at 20 °C for 15 min to form a yellow solution, presumably containing the Cp₂Fe byproduct, that was removed by filtration. The red solids that remained atop the frit were washed with pentane (5 mL × 2) until the washings were colorless, and dried in vacuo. Crystallization from benzene/THF gave bright orange microcrystals of [Cp₂Fe]OTf (37.0 mg, 68% yield). 1H NMR (282 MHz, CDCl₃, 20 °C): δ = –76.9 ppm. Anal. Calcd for H₂, by AgOTf. 19F NMR (376 MHz, CDCl₃, 20 °C): –144 (sept, JFP = 20.1 Hz) ppm. 31P NMR (162 MHz, CDCl₃, 20 °C): δ = –144 (sept, JFP = 715 Hz) ppm.

**Oxidative Cleavage of μ-(PhCH(NN)₂)Mo[N=0-Bu]Ar₂]-[H]₂ by AgOTf.** The [H]-I (45.1 mg, 0.0304 mmol) was dissolved in Et₂O (20 mL). To the solution was added AgOTf (15.6 mg, 0.0607 mmol) and the mixture was stirred for 2 h at room temperature. The mixture was filtered and the volatiles in the filtrate were removed in vacuo. Crystallization from benzene/THF layer with n-pentane gave bright orange microcrystals of [PhCH(NN)₂]Mo[N=0-Bu]Ar₂]-[AgOTf] (37.0 mg, 68% yield). 1H NMR data is identical to that reported for [H]-I][AlCl₄]. 19F NMR (376 MHz, CDCl₃, 20 °C): δ = –73 (d, 1JFP = 715 Hz) ppm. 31P NMR (162 MHz, CDCl₃, 20 °C): δ = –144 (sept, 1JFP = 715 Hz) ppm.

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**Supporting Information Available:** ESI-MS, 1H NMR, IR, Raman and CV data for compounds [1-R][AlCl₄] in addition to computational details are included in the supporting information. This material is available free of charge via the Internet at http://pubs.acs.org.

**References:**

Due to the strong kinematic coupling between oscillators in [HAI][Cl], the origin of the IR- and Raman-active band found at 1173 cm⁻¹ is uncertain.

A Flack x parameter of -0.050(27) was calculated by SHELXL.


The preparation of a series of cationic diazooalkane complexes \([4-\text{RC}_6\text{H}_4\text{C(H)NNMo}[\text{t-BuAr}_3]][\text{AlCl}_4]\), \([1-\text{R}][\text{AlCl}_4]\) (\(\text{R} = \text{NMe}_2, \text{Me}, \text{H}, \text{Br}, \text{CN}, \text{Ar} = 3,5-\text{C}_6\text{H}_3\text{Me}_2\)) is described, and the spectroscopic properties of these complexes are compared. One-electron reduction of \([1-\text{R}][\text{AlCl}_4]\) forms the \(\text{C}\text{–C}\) bonded dimer, \(\mu-(4-\text{RC}_6\text{H}_4\text{C(H)NN})_2[\text{Mo}(\text{t-BuAr}_3)]_2\). The \(\text{C}\text{–C}\) bond in \([1-\text{R}]_2\) is redox non-innocent and is broken upon oxidation to regenerate \([1-\text{R}]\).