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### **EPR/ENDOR and theoretical study of the Jahn-Teller active [HIPTN3N]Mo(V)L complexes (L = N−, NH)**

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#### **Abstract**

The molybdenum trisamidoamine (TAA) complex [**Mo**] (=(3,5-(2,4,6-i-

 $Pr_3C_6H_2$ )<sub>2</sub>C<sub>6</sub>H<sub>3</sub>NCH<sub>2</sub>CH<sub>2</sub>N)Mo) carries out catalytic reduction of N<sub>2</sub> to ammonia by protons and electrons at room temperature. A key intermediate in the proposed [**Mo**] nitrogen reduction cycle is nitrido-Mo(VI),  $[Mo(VI)]N$ : the addition of  $[e^{-}/H^{+}]$  to  $[Mo(VI)]N$  to generate  $[Mo(V)]NH$ might in principle follow one of three possible pathways: direct proton-coupled electron transfer; H+ first, then e−; e− then H+. In this study, the paramagnetic Mo(V) intermediate {[**Mo**]N}− and [**Mo**]NH transfer product were generated by irradiating the diamagnetic [**Mo**]N, {[**Mo**]NH}<sup>+</sup> Mo(VI) complexes respectively, with  $\gamma$ -rays at 77 K, and their electronic and geometric structures were characterized by electron paramagnetic resonance (EPR), electron nuclear double resonance (ENDOR) spectroscopies combined with quantum chemical computations. In combination with previous X-ray studies this creates the rare situation where each one of the four possible states of an [e−/H+] delivery has been characterized. Because of the degeneracy of the electronic ground states of both, {[**Mo(V)**]N}− and [**Mo(V)**]NH, only multi-reference based methods such as the complete active space self-consistent field (CASSCF) and related methods provide a qualitatively correct description of the electronic ground state and vibronic coupling. The molecular g-values of {[**Mo**]N}− and [**Mo**]NH exhibit large deviations from the free electron value g<sup>e</sup> . Their actual values reflect the relative strengths of vibronic and spin-orbit coupling. In the course of the computational treatment, the utility and limitations of a formal two-state model that describes this competition between couplings are illustrated, and implications of our results for the chemical reactivity of these states are discussed.

δCurrent Address: School of Mathematics and Physical Sciences, University of Hull, Cottingham Road, HU6 7RX, United Kingdom **Supporting Information Available:** Orientation selective<sup>14,15</sup>N, <sup>1</sup>H ENDOR spectra of Mo<sup>14,15</sup>N<sup>−</sup>, Mo<sup>14,15</sup>N<sup>1</sup>H; ENDOR simulations; calculated active orbitals of MoN−, MoNH; Townes-Dailey Quadrupole Analysis; 12 figures in all.

#### **TOC image**



#### **Introduction**

The molybdenum trisamidoamine (TAA) complex [**Mo**] (=(3,5-(2,4,6-i- $Pr_3C_6H_2$ )<sub>2</sub>C<sub>6</sub>H<sub>3</sub>NCH<sub>2</sub>CH<sub>2</sub>N)M<sub>o</sub>) shown in chart 1 is the active species in the catalytic reduction of  $N_2$  to ammonia by protons and electrons at room temperature.<sup>1–3</sup> Despite its low turnover rate this complex represented a major advance in the field of catalytic nitrogen fixation and activation, as the first well-defined molecular catalyst of nitrogen reduction. Some of the key intermediates of the reaction cycle have been crystallized and studied spectroscopically,<sup>1–4</sup> the magnetic properties of the key intermediates,  $[Mo]N_2$  and [Mo]NH<sub>3</sub>, as well as of the CO inhibited form, have been studied in detail,<sup>5,6</sup> and the entire reaction pathway has been investigated with quantum chemical methods.<sup>7-11</sup>

The  $[Mo]N_2$ ,  $[Mo]NH_3$ , and  $[Mo]CO$  complexes are low spin  $(S = \frac{1}{2}) d^3$ , which is rare in metal coordination compounds. It has been demonstrated that their unique geometric and electronic structure have a remarkable effect on the molecular g-tensor of the three complexes. The TAA ligand enforces a trigonal symmetry on the molybdenum center leading to partial occupation of a set of doubly occupied orbitals. This partial occupation in turn leads to strong 'in-state' spin-orbit coupling (SOC) and Jahn-Teller (JT) vibronic coupling that compete with each other, resulting in the so-called 'Pseudo-Jahn-Teller' (PJT) effect.12 The combined results of ligand field theory analysis and quantum chemical methods revealed that the molecular g-tensor is governed by the 'in-state' SOC, which is effectively diminished with increasing JT distortion. Hence the g-values can be regarded as sensitive probe of the ratio between SOC and JT coupling. More recently, the TAA ligand has received attention owing to its use in iron-based single molecule magnets  $(SMM)$ ,  $^{13,14}$ and parallel efforts in the synthesis and characterization of trigonal Fe complexes,  $15-20$ likewise reveal such properties, and indeed have yielded nitrogen reduction catalysts.<sup>21,22</sup>

In the proposed [Mo] nitrogen reduction cycle there are four Mo(V) paramagnetic  $(S = \frac{1}{2})$ intermediates, namely  $[Mo]N_2H_2$ ,  $\{[Mo]N_2H_3\}^+$ ,  $[Mo]NH$ , and  $\{[Mo]NH_2\}^+$ , in addition to the two  $Mo(III)$  complexes.<sup>1–3</sup> Though the diamagnetic  $Mo(VI)$  intermediates {[**Mo**]N2H2} <sup>+</sup>, [**Mo**]N, and {[**Mo**]NH}+ that fall just one step before the Mo(V) species in the catalytic cycle are well characterized, the Mo(V) intermediates could not be observed ([**Mo**]N2H2) and/or isolated ([**Mo**]NH) during synthesis.<sup>4</sup> As illustrated in Figure 1, the addition of [e−/H+] to [**Mo(VI)**]N to generate [**Mo(V)**]NH might in principle follow one of three possible pathways: direct proton-couple ET; H<sup>+</sup> first, then  $e^-$ ;  $e^-$  then H<sup>+.</sup> In the

present study, the paramagnetic Mo(V) species {[**Mo**]N}−, and [**Mo**]NH were instead generated by irradiating the diamagnetic [**Mo**]N, {[**Mo**]NH}+ complexes respectively, with γ-rays at 77 K. Although cyclic voltammetry shows that {[**Mo**]N}− is formed reversibly at low potential ( $E_{1/2}$  = −2.68 V *vs.* Fc<sup>+</sup>/Fc (Pt)),<sup>23</sup> it has never been isolated. Thus, the combination of the previous X-ray diffraction studies and the present spectroscopic/ theoretical investigation of {[**Mo(V)**]N}− and {[**Mo**]NH} enable the rare situation where each one of the four possible states of [e<sup>−</sup>/H<sup>+</sup>] delivery can be characterized.

We here use the electron paramagnetic resonance (EPR), electron nuclear double resonance (ENDOR) spectroscopic techniques combined with the computational studies to characterize the electronic and geometric structure of these two Mo(V) intermediates,{[**Mo**]N}−, [Mo]NH. The first part of this report describes the investigation of the electronic structures of cryoreduced {[**Mo**]N}− and the protonated form [**Mo**]NH with quantum chemical methods. The degeneracy of the electronic ground states of both compounds (vide infra) prohibits the usage of density functional theory (DFT) for the calculation of the electronic structure and magnetic properties. Only multi-reference based methods such as the complete active space self-consistent field (CASSCF) and related methods provide the necessary flexibility for a qualitatively correct description of the electronic ground state and vibronic coupling. Moreover the strong 'in-state' SOC cannot be treated by regular linear response approaches but requires an explicit evaluation via quasi-degenerate perturbation theory. In the second part the EPR spectra of the two complexes are reported and first interpreted in terms of a two-orbital model for a doubly degenerate  ${}^{2}E$  ground state, then analyzed in the light of the quantum-chemical results of the first part. The molecular g-values of {[**Mo**]N}<sup>−</sup> and [**Mo**]NH exhibit large deviations from the free electron value g<sup>e</sup> . Their actual values serve as a probe of the relative strengths of vibronic and spin-orbit coupling. The third and final part discusses bonding to the ligands surrounding Mo as revealed by their nuclear hyperfine interactions.  $^{14,15}N$  ENDOR spectroscopy identifies the axial nitride ligand of {[**Mo**]N}−, and shows it to have a small isotropic nuclear hyperfine coupling (∼ 3 MHz), a result nicely captured by the calculations. In contrast, the calculations predicts a large isotropic hyperfine coupling (∼ 10 MHz) for the axial imido nitrogen of **Mo**NH, and rationalize an inability to measure these couplings directly.  $1.2$  H ENDOR experiments agree with the calculation of a bent structure for Mo-N-H with a substantial isotropic coupling for the imido proton. In the course of this treatment, the utility and limitations of the two-state model are illustrated, and implications for the chemical reactivity of these states are discussed.

#### **Materials and Methods**

#### **Synthesis**

Solutions of [Mo]N, and  $\{[Mo]NH\}$ <sup>+</sup> were synthesized as described in the literature.<sup>4</sup> The solutions were frozen in liquid nitrogen for cryoreduction and for subsequent EPR/ENDOR measurements. Solutions of diamagnetic [**Mo**]N, and {Mo(VI)NH}+ were cryoreduced at 77 K, using γ-rays to form paramagnetic {[**Mo**]N}−, and [**Mo**]NH respectively<sup>24</sup> .

#### **EPR and ENDOR Measurements**

The 35 GHz pulsed EPR and ENDOR measurements at ∼2 K, were carried out using a locally constructed spectrometer described in detail previously<sup>25</sup>. Field swept electron spin echo (ESE) EPR were collected with the two pulse echo sequence:  $π/2$ - $τ$ - $π$ - $recho$ . Pulsed ENDOR experiments employed either the Davies  $(\pi$ -T- $\pi/2$ - $\tau$ - $\pi$ - $\tau$ - $echo$ ), or the MIMS sequence  $(π/2-τ-π/2-T-π/2-τ-echo)$ , where T is the time interval for which the RF pulse is applied<sup>26</sup>. Data acquisition for the pulse measurements utilized the SpecMan software package27 ([http://specman.4epr.com\)](http://specman.4epr.com) in conjunction with a Spin-Core PulseBlaster ESR\_PRO 400 MHz word generator and an Agilent technologies Acquiris DP235 500 MS/sec digitizer.

For a single molecular orientation and for nuclei with a nuclear spin of  $I = \frac{1}{2}$  ( ${}^{1}H$ ,  ${}^{15}N$ ), the ENDOR transitions for the  $m_s = \pm \frac{1}{2}$  electron manifolds are observed at frequencies given by the equation

$$
\nu_{\pm} = |\nu_{\rm n} \pm A/2| \quad (1)
$$

where  $v_n$  is the nuclear Larmor frequency and A is the orientation-dependent hyperfine coupling. For I  $\pm 1$  (<sup>14</sup>N, <sup>2</sup>H), the two ENDOR lines are further split by the orientationdependent nuclear quadrupole interaction (3P). The peaks are then observed at frequencies dictated by a modified version of the equation above

$$
\nu_{\pm,\pm} = |\nu_{\pm} \pm 3P/2|
$$
 (2)

It is well documented that by analyzing the 2D field-frequency ENDOR spectra collected at various fields across the EPR envelope, the complete nuclear hyperfine and quadrupole principal values could be determined.28–31

#### **Computational Details**

All calculations on Mo(V) complexes performed in this work were conducted with the ORCA quantum chemistry program package.<sup>32</sup> Density functional theory as well *ab initio* wave function calculations used the zero'th order regular approximation (ZORA) utilizing a model potential introduced by van Wüllen.33,34 Accordingly, the segmented all-electron relativistically contraction (SARC) versions of the TZVP and SV(P) basis sets, SARC-TZVP and SARC-SV(P), were used throughout the work.<sup>35</sup> The Resolution of Identity (RI) approximation was applied for all integral transformations.

Analogous to a previous investigation of the  $[Mo]$  complex,<sup>5</sup> a slightly reduced model structure of the full complex geometry has been used for many calculations. The applied model is in accordance with model 1″ of reference 8. It features the molybdenum center, the nitrido/imido ligand and the TAA ligand but the large HIPT groups have been replaced by ethyl groups in order to safe substantial computational costs without completely losing the electron donating effect of the HIPT groups (Figure 2). Geometry optimization of the model structure comprised of two steps. First, the full complex geometry has been fully optimized

using the BP86 $36,37$  functional together with the SARC-TZVP (Mo, N) and the SARC- $SV(P)$  (C, H) basis set.<sup>35</sup> Then the HIPT groups have been replaced by ethyl groups, which then were optimized separately at the same level of theory. Furthermore a second set of structures was generated by optimizing the model structure with respect to the average of the first two electronic states (the two members of  ${}^{2}E$ -term) utilizing the state-averaged CAS-SCF(7,8) method (vide infra). In the following these structures will be referred to as 'symmetric' structures since their first coordination shell exhibits almost perfect  $C_{3V}$ symmetry. The most important structural parameters of the first coordination shell of the fully relaxed and the symmetric structures can be found in the Supporting Information. In the case of the optimization to the symmetric limit of [**Mo**]NH the Mo-Nimido-H bond angle is close to 180° which is an artifact of the energy minimization with respect to the average of the lowest two states. Of course, this chemically unreasonable geometry leads to a significant change of the metal ligand bond pattern. Therefore we have manually adjusted this angle to a value of 154° (the value obtained from the fully relaxed optimization) for the multiplet calculations in the symmetric limit (vide infra).

In the course of this work three sets of multireference ab initio calculations were conducted that employ the CASSCF method with different active spaces and varying levels of treatment of dynamic electron correlation effects. The first set of calculations consists of the aforementioned geometry optimizations to the symmetric limit and a set of scans of the Jahn-Teller potential energy surface. In both cases an active space with 7 electrons in 8 orbitals has been chosen, with the shape of the active orbitals being the result of an optimization according to the variational principle. For {[**Mo**]N}− the active orbitals correspond to the five metal d orbitals as well as the three p-orbitals of the nitrido axial ligand (Figure 3). By contrast, the active orbitals of [**Mo**]NH comprise one amido p orbital, three metal d orbitals ( $d_{xz}$ ,  $d_{yz}$  and a combination of the  $d_{xy}$  and  $d_{x^2-y^2}$  orbitals) as well as four p-orbitals ( $2p_{x/y}$  and  $3p_{x/y}$ ) of the axial nitrido ligand. These choices are expected to provide a balanced description of the Mo-N bond in both complexes, and hence, is a good basis for their qualitative discussion.

Of course, the shape and composition of the active orbitals alters as the corresponding geometry changes but their main character remained unchanged throughout. Dynamical electron correlation was taken into account in the framework of the second-order N-electron valence perturbation theory (NEVPT2) scheme for the potential energy surface scans while they were neglected during the geometry optimizations.

The second set of calculations comprises only single-point energy and g-tensor evaluations. They feature an active space of 13 electrons in 11 orbitals and use NEVPT2 to incorporate dynamical electron correlation. In the case of [**Mo**]N− the 11 active orbitals that resulted from optimizations with respect to the variational principle include the five metal d-based orbitals, the three p-orbitals and the one s-orbital of the nitrido axial ligand and two σbonding orbitals associated with the equatorial amido ligands (cf Figure SI 8). For [**Mo**]NH the active orbitals correspond to four metal d orbitals ( $d_{xz}$ ,  $d_{yz}$ ,  $d_{xy}$  and  $d_{x^2-y^2}$ ), four porbitals of the axial ligand and three  $\sigma$ -bonding orbitals associated with the equatorial amido ligands (cf. Figure SI 9), again the result of optimizations with respect to the variational

principle. This second set of calculations is the most elaborate and also most accurate calculation performed. The inclusion of the equatorial ligand bonding orbitals in the active space will ensure that the ligand field induced by the N/NH as well as the supporting ligand is balanced.

A third set of calculations was performed that only features the minimal active space of 1 electron in the 5 metal d-based orbitals (cf. Figure SI 10 and Figure SI 11). For this set of calculations, dynamical electron correlation effects are described by means of the difference dedicated configuration interaction with a maximum number of two degrees of freedom (DDCI2). In the course of the DDCI2 procedure thresholds of  $T_{pre} = 10^{-4}$  for the selection of configurations in the reference space and  $T_{\text{sel}} = 10^{-6}$  for the selection of configurations in the CI part were applied. The higher order correlation terms included in DDCI2 make up for some of the deficiencies of the small active spaces and are a convenient basis for discussing some qualitative features of the investigated systems.

Note, that in all three sets of calculations multiple electronic states were evaluated and that the CASSCF orbitals were optimized in a state-averaged fashion. The actual number of calculated states will be indicated in the description of each individual calculation.

Molecular g-tensors have been calculated using either linear response DFT or, for the multireference calculations, the formalism introduced by Gerloch and McMeeking.<sup>38,39</sup> In order to obtain g-tensors from linear response DFT, a set of coupled perturbed Kohn-Sham equations had to be solved for a set of self-consistent Kohn-Sham orbitals and a magnetic field perturbation. A more detailed description of this procedure can be found in reference<sup>38</sup>. The formalism proposed by Gerloch and McMeeking was used for g-tensor calculations with multireference wave functions: Complete Active Space Self-Consistent Field (m,n) calculations  $(CASSCF(m,n))$  where m is the number of active electrons in n active orbitals were conducted at the optimized geometries. Dynamic correlation effects were then taken into account by either the NEVPT2 or the DDCI2 scheme. $40-44$  In the following we will denote a set of CASSCF(m,n) and subsequent NEVPT2 or DCCI2 calculations as NEVPT2(m,n) or DCCI2(m,n). However, it should be noted that in the case of NEVPT2 only the energy correction to the state energies from perturbation theory is used whereas the form of the wave function is left unchanged. The resulting set of eigenstates  $\{|\vec{l}\rangle\}$  forms a basis in which the Born-Oppenheimer Hamiltonian is presumed to be diagonal. On top of these non-relativistic calculations, SOC is introduced in the framework of quasi-degenerate perturbation theory (QDPT). This means that the SOC operator is diagonalized in the basis of eigenstates  $\{|\vec{l}\rangle\}$ . For the evaluation of SOC integrals the spin-orbit mean-field Hamiltonian  $\hat{H}_{SOMF}$  is used, which appears as an effective one-electron operator<sup>45,46</sup>:

$$
\hat{H}_{SOMF} = \sum_{i} \hat{h}^{SOC}(i)\hat{s}(i)
$$
\n(3)

The presented approach accounts for SOC to infinite order but suffers from truncation of the incomplete basis of  $\{|\Lambda\rangle\}$  functions. The diagonalization yields eigenfunctions which appear as pairwise degenerate functions  $\phi$  and  $\overline{\phi}$ , the so-called Kramer's pairs. An external

magnetic field splits the degeneracy of these pairs via the Zeeman term  $\hat{H}_{Zee}$  that is then added to the Hamiltonian:

$$
\dot{H} = H_{BO} + \dot{H}_{SOMF} + \dot{H}_{Zee} \quad (4)
$$

with

$$
\hat{H}_{\text{Zee}} = \mu_B \left( \hat{L} + g_e \hat{S} \right) \boldsymbol{B} \tag{5}
$$

Here,  $\mu_B$  is the Bohr magneton,  $g_e$  is the Landé factor, **B** is the magnetic flux density and and  $\hat{S}$  are the total orbital and spin angular momentum operators, respectively. It has been shown that diagonalization of the Zeeman term in the basis of the Kramer's doublet is equivalent to finding the eigenvalues of the Zeeman term in the basis of fictitious spin functions  $|+\rangle$  and  $|-\rangle$  in the spin-Hamiltonian. Hence, the square of the g-tensors can be calculated according to<sup>39,47</sup> ( $\mu$ ,  $\nu=x, y, z$ ):

$$
\Gamma_{\mu\nu} = \left(gg^T\right)_{\mu\nu} = 2 \sum_{I,J = \phi,\overline{\phi}} \left\langle I \left| \hat{L}_{\mu} + g_e S_{\mu} \right| J \right\rangle \left\langle J \left| \hat{L}_{\nu} + g_e S_{\nu} \right| I \right\rangle \tag{6}
$$

#### **Results**

#### **Electronic structure and PJT**

The tripodal ligand enforces an idealized  $C_{3V}$  symmetric coordination environment to the central Mo(V) ion. As outlined previously, it is this special environment that accounts for the unusual magnetic properties of the derivatives of the  $[Mo]$  complex.<sup>5</sup> In a  $C_{3V}$ -symmetric ligand field the five metal d-orbitals are split into two degenerate pairs (e) and one nondegenerate orbital  $(a_1)$ , where the orbitals can be expressed either as conventional, real dorbitals or as eigenfunctions of the z-component of the orbital angular momentum,  $m_l = \pm 2$ ,  $\pm 1$ , 0 (l=2):

$$
3e = [d_{xz}, d_{yz}] = \left[ \begin{array}{c} | \ 1 \end{array}, \left| -1 \right\rangle \right]
$$

$$
4e = \begin{bmatrix} d_{x^2-y^2}, d_{yz} \end{bmatrix} = \begin{bmatrix} |2\rangle, |-2\rangle \end{bmatrix}
$$
 (7)

$$
4a_1 = [d_{z^2}] = [ |0\rangle ]
$$

Hence, the d<sup>1</sup> configuration of the {[Mo]N}<sup>−</sup> and [Mo]NH complexes gives rise to three different doublet states: Two orbitally doubly degenerate <sup>2</sup>E states and one orbitally non-

degenerate  ${}^{2}A_1$  state. NEVPT2(13,11) calculations including the 15 lowest states on the 'symmetric' structures of both complexes allowed for a qualitative and quantitative analysis of their d-d multiplet structure. The corresponding molecular orbitals involved in these calculations of course are linear combinations of these metal d-orbitals and ligand orbitals. Table 3 summarizes the calculated energetic levels of the d-multiplets and their respective dominant configurations as obtained from the CASSCF calculation.

The energies presented in Table 3 show that addition of a proton to {[**Mo**]N}− leads to a change in the electronic ground state configuration. In the case of  $\{[\text{Mo}]\text{N}\}^-$  the 4e(x<sup>2</sup>y 2 ,xy) orbital manifold is singly occupied whereas the electronic ground state of [**Mo**]NH corresponds to a singly-occupied 3e(xz,yz) manifold. For both complexes the complementary <sup>2</sup>E state is the first excited state while the respective  ${}^2A_1$  state is highest in energy. Due to the size of the active space and the limited number of calculated states, the actual energy of the  ${}^{2}A_1$  state could not be determined since a series of charge transfer states occur lower in energy than the highest d-d state. In the case of [**Mo**]NH the orbital optimization even rotated the metal  $d_z$  orbital out of the active space which impedes a calculation of the  ${}^2A_1$  state. Consequently, in Table 3 the excitation energy for the  ${}^2A_1$  states obtained from the corresponding NEVPT2(1,5) are given for this state. They can be regarded as reasonable approximations to the results that would have been obtained from NEVPT2(13,11) calculations. For the present case of  $d<sup>1</sup>$  configurations the qualitative frontier orbital schemes highlighted in Figure 4 directly correspond to the ordering of the calculated d-d excited states. They differ in the energetic order of the 3e and 4e orbitals. A qualitative explanation for these results can be given in the framework of ligand field theory. The ligand field created by the TAA ligand has a dominating component perpendicular to the molecular axis and arises from the Mo-N<sub>amido</sub>  $\sigma$ -interactions. It destabilizes the 4e(x<sup>2</sup> $y^2$ ,xy) orbitals relative to the 3e(xz,yz) orbitals. On the other hand, the axial ligand generates a ligand field along the molecular axis that destabilizes the metal d-orbitals in the order  $d_z^2$  >  $d_{\text{XZ, yZ}} > d_{\text{XY, x}}^2 - y^2$ . Accordingly the <sup>2</sup>A<sub>1</sub> state is by far the highest in energy among the d-d excited states for both complexes.

The energetic order of the 3e and 4e orbitals depends on the relative strengths of ligand fields created by the TAA ligand and the axial ligand. Since the TAA ligand is a common feature of both complexes, the energetic order depends only on the ligand field strength of the axial ligand. In previous works it has been found that for  $[Mo]L$  ( $L = NH<sub>3</sub>$ ,  $N<sub>2</sub>$ , CO) the 3e orbitals are lower in energy than the 4e orbitals. The same situation is met for [**Mo**]NH. In case of {[**Mo**]N}− due to the extraordinarily strong π-interaction created by the nitrido ligand, the energetic order of 3e and 4e orbitals is inversed. Accordingly the numbering used here no longer refers to the energetic order of the orbitals. All interactions between the ligand and metal orbitals are summarized in the qualitative molecular orbital scheme presented in Figure 4.

Despite their different electronic ground state configurations {[**Mo**]N}−, and [**Mo**]NH share some critical properties. They both have a doubly degenerate  ${}^{2}E$  ground state that is subject to JT distortion which can be described by the well-known  $E \otimes e$  coupling scheme.<sup>12</sup> Furthermore the  ${}^{2}E$  ground state exhibits unquenched orbital angular momentum leading to strong 'in-state' spin-orbit coupling (SOC). The SOC splits the orbital and spin-degeneracy

and thus competes with the JT distortion resulting in the so-called 'Pseudo-Jahn-Tellereffect'. As described in detail elsewhere,<sup>5</sup> the PJT effect forces the system to distort along a 'composite' mode of vibration leading to two separate molecular adiabatic potential energy surfaces (APES) with energies defined in terms of the force constant,  $K$ , of the "composite" mode along which the distortion  $\rho$  occurs,  $\lambda$  the SOC constant and F, the linear Jahn-Teller coupling constant.

$$
W_{\pm} = \frac{K\rho^2}{2} \pm \sqrt{\lambda^2 + (2F\rho)^2}
$$
 (8a)

The equilibrium displacement on the lower surface occurs at a distortion,  $\rho_0$ , at which point the

$$
\rho_0 = \sqrt{\left(\frac{F}{K}\right)^2 - \left(\frac{\lambda}{2F}\right)^2}
$$
 (8b)

By comparison of the optimized geometries of the symmetric and the fully optimized structures it was possible to determine the nature of the mode of distortion. As in the previous ([Mo]L,  $L = N_2$ , NH<sub>3</sub> and CO) cases they are dominated by transversal movement of the axial ligands and bending motion of the equatorial ligands as illustrated in Figure 5.

#### **Magnetic Measurements**

**EPR g-values and the formal PJT theory—**Figure 6 shows the 35 GHz, EPR spectra of the frozen solution of [Mo]NH, and {[Mo]N}− recorded at 2K. The EPR spectrum of {[Mo]N}− shows axial symmetry, g<sup>ǁ</sup> = 1.69 < g⊥ = 1.91. The g tensor for [**Mo**]NH is nearaxial,  $g = [(g_{\parallel} = g_3) = 1.86 < {g_1 = 1.98, g_2 = 1.96}]$ , with a smaller g-spread than {[**Mo**]N}−. The similarity of the EPR spectra for the two complexes despite the fact that they have different  ${}^{2}E$  ground states can be understood in terms of the 'pseudo Jahn-Teller' (PJT) effect wherein spin-orbit coupling competes with interactions that split the orbital degeneracy, lowers the molecular symmetry, and lowers the total energy of the molecule.<sup>5</sup> Foremost among the degeneracy-lifting terms is the JT vibronic coupling to a symmetrylowering distortion along a composite 'interaction'  $e_a$  coordinate, but deviations from trigonal symmetry such as occurs in [Mo]NH, other electronic interactions, and indeed noncovalent solvent effects in solution or crystal packing effects in a crystal, can contribute.

Within the previously introduced two-orbital model for a doubly degenerate <sup>2</sup>E ground state, the mixing and splitting of the orbital doublet can be described formally<sup>5</sup> in terms of a dimensionless parameter,  $r$ , and fictitious angle  $\theta$ 

$$
\tan 2\theta \equiv r = \frac{2V}{\lambda} \lambda = k\lambda_{\text{soc}}
$$
 (9)

The numerator is a sum of two types of terms,  $V = V_{vib} + V_l$ . The first term is the vibronic coupling energy at the equilibrium distorted position on the ground APES,  $V_{vib} = F\rho_0$ , The second term in the numerator,  $V<sub>k</sub>$  is the sum of the additional contributions enumerated in the previous paragraph. The denominator,  $\lambda$  is the spin-orbit coupling (SOC) parameter, which traditionally is taken to be the ionic SOC constant,  $\lambda_{SOC}$ , multiplied by a 'covalency parameter',  $k$ , defined such that  $(1-k)$  represents the fraction of d-electron density delocalized onto the ligands. Within this model the g values can be written  $as<sup>5</sup>$ 

$$
[2e^1]g_{\parallel} = 2(1 - m_l k \cos 2\theta) \quad (10a)
$$

$$
[2e^1]g_{\perp} = 2\sin 2\theta \quad (10b)
$$

and the The PJT splitting between the ground APES minimum and the excited APES at the equilibrium value of the distortion,

$$
E_{JT} = \lambda \sqrt{1 + \tan(2\theta)^2}
$$
 (11)

According to the quantum calculations presented above,  $m_l = 2$  for  $\{[\text{Mo}]\text{N}\}^-$ , but  $m_l = 1$  for [Mo]NH, as is the case for the d<sup>3</sup> Mo(III) complexes previously studied by paramagnetic resonance methods.5,6

The physical picture behind the deviation of the g  $(g_{\parallel, g_{\perp}})$  values from the free electron value can be understood as follows. In the limit of non-distorted trigonal symmetry (no JT effect,  $2\theta = 0$ ), the strong 'in-state' SOC leads to antiferromagnetic coupling of orbital and spin angular momentum along the z axis resulting in a  $g_{\parallel}$  value of -2 and 0 for  $\{[\text{Mo}]N\}^{-}$  (4e)<sup>1</sup> and  $[Mo]NH (3e)^1$ , respectively, in the ionic limit of  $k = 1$ . As the signs of the g-values are not resolved in ordinary EPR measurements, one would measure a value of  $|g_{\parallel}| = +2$  for {[**Mo**]N}− in a three-fold symmetric conformation. As shown in Figure 7, the absolute value of g|| for {[**Mo**]N}− rapidly decreases from 2 to 0 and then slowly increases towards the free electron value g<sup>e</sup> with increasing Jahn Teller splitting. For [**Mo**]NH g|| starts at 0 in the limit of no distortion and increases as the PJT becomes larger.

As a secondary effect, the SOC along the z-axis quenches the magnetic moment perpendicular to it. Thus  $g_{\perp}$  vanishes in the limit of a three-fold symmetry,  $r = 0$ . The Jahn-Teller splitting progressively diminishes the orbital angular momentum which in turn reduces the 'in-state' SOC. This is reflected in a progressive increase of g⊥ towards the freespin value,  $g_e = 2$  as presented in Figure 7.

Treating the measured g-values with **Eqs 9, 10** gives a small value,  $r = 3.2$  ( $E_{JT} = 3352$ cm<sup>-1</sup>) for {[**Mo**]N}<sup>-</sup> and a larger value for [**Mo**]NH,  $r = 7$  (E<sub>IT</sub> = 7071 cm<sup>-1</sup>). The larger r for Mo-N-H can be attributed to the non-linear geometry, which, along with the larger SOC parameter for Mo(V) (1000cm<sup>-1</sup>) compared to Mo(III) (800 cm<sup>-1</sup>) acts to enhance the

splitting of the orbital degeneracy. The rhombicity of the g-tensor of [**Mo**]NH reflects SOC to other d orbital, that is not included in the two-orbital model.

For {[**Mo**]N}−, this treatment gives unrealistically low values for the 'covalency parameter', <sup>k</sup>, which commonly are interpreted as representing the metal-d character of the 3/4e orbital,  $k = 0.25$ . In part, a low value can be understood by noting that in the idealized  $C_{3v}$  symmetry of  $\{[\text{Mo}]\}$ <sup>-</sup> Telser and McGarvey showed how this symmetry-lowering mixes the  $e_b 3E$ and 4E orbitals,<sup>48</sup> which diminishes the covalency parameter,  $k$ ,<sup>18</sup> for  $\{[\text{Mo}]N\}$ <sup>-</sup>. However, it would seem that this low value also reflects limitations to the two-orbital model. In the case of  $[Mo]NH$  a more realistic value of  $k = 0.5$  is obtained. A discussion of covalency and the mixing of states with different parentage from the point of view of quantum chemical calculations for both complexes will be given below.

#### **Quantum Chemical Computations of g-values**

**[Mo]NH:** The formal PJT analysis predicts a large splitting between the equilibrium ground state APES and the first excited APES for [**Mo**]NH. A NEVPT2(13,11) calculation of the lowest ten electronic states on the model structure of the complex (Figure 2) supports this assessment. It yielded an excitation energy of  $6000 \text{ cm}^{-1}$  for the first excited state, which is on the same order as the value obtained from the two-orbital model. This large intra- ${}^{2}E$ splitting is a result of the presence of the imido proton that is effective in breaking the symmetry. Evaluation of the g-tensor at the same level of theory yields values of  $g_1 = 1.99$ ,  $g_2 = 1.96$  and  $g_3 = 1.87$  which is in excellent agreement with the experimental values and consistent with the results obtained from DDCI2(1,5) ( $g_1 = 1.98$ ,  $g_2 = 1.95$  and  $g_3 = 1.80$ ). Considering the large PJT splitting,  $E_{JT}$ , the electronic ground state of the desymmetrized structure may be regarded as genuinely non-degenerate. Thus, the application of the standard linear response DFT methodology to the calculation of the molecular g-tensor is reasonable. A great advantage of standard DFT calculations is their low computational cost, which allows the inclusion of the full HIPT groups in the g-tensor calculation. At this point it has to be noted that linear response DFT incorporates SOC only to second order in perturbation theory while the NEV-PT2 approach treats SOC to infinite order. However, the g-values obtained from linear response DFT ( $g_1 = 1.99$ ,  $g_2 = 1.96$ ,  $g_3 = 1.88$ ) are almost equally close to the experimental data as the above presented NEVPT2 values. In order to elucidate the effect of truncation of the HIPT groups on the g-tensor a linear response DFT calculation was also performed for the model structure of [**Mo**]NH. It gives almost identical results (g<sub>1</sub>  $= 1.99$ ,  $g_2 = 1.97$ ,  $g_3 = 1.88$ ) as the calculation on the full complex. This indicates that truncation of the model does not significantly influence the results of NEVPT2 g-tensor calculations.

The average metal-d character of the 3e orbitals in the Kohn-Sham determinant is 47%, which is close to the value obtained from the two-orbital model. This corresponds to strongly covalent bonds between the metal and the ligands, especially the axial imido ligand, although one has to keep in mind that DFT tends to overestimate covalency in transition metal complexes.<sup>49</sup> In the large active-space CASSCF(13,11) calculations, which should provide a rather balanced description of the [**Mo**]NH bonds, the metal character in the openshell orbitals is 59% and 69% respectively. For comparison, the averaged metal-d character

of the natural DDCI2 orbitals that correspond to the 3e manifold in [**Mo**]NH is 55% which is close to the value obtained for [**Mo**]NH in the two-orbital model. These numbers appear to be sensible and the description of the bond is further supported by the good agreement between calculated and experimental g-values that react sensitively to both the intra-  ${}^{2}E$ excitation energy as well as the covalent dilution of the d-orbitals.

An analysis of the spin distribution evaluated at the DDCI2(1,5) level provides insight into the electronic structure at the distorted geometry of [**Mo**]NH. Figure 8 shows the spin density at an isosurface value of 0.003 unpaired electrons. In agreement with the CASSCF results, it reveals that the electron spin is mainly located in an antibonding combination of Mo  $d_{xz}/d_{yz}$  orbital and imido  $p_x/p_y$  orbital. This finding supports the assessments made above about the electronic structure of [**Mo**]NH (Figure 4). Furthermore it is seen that the spin distribution is significantly tilted with respect to the molecular z-axis resulting from the geometrical distortion and leading to a non-negligible admixture of Mo  $d_{x^2-y^2}/d_{xy}$  and equatorial ligand p orbitals. In a configuration interaction (CI) picture this spin distribution corresponds to a distortion-induced mixing of  ${}^2\!E_b$  states to the electronic ground  ${}^2\!E_a$  state.

**{[Mo]N}−:** The two-orbital model predicts a significantly smaller PJT for {[**Mo**]N}<sup>−</sup> compared to [**Mo**]NH (vide supra). The NEVPT2(13,11) calculation for the lowest five states of the model structure of {[**Mo**]N}− supports this result. It yielded an energy difference of 4200 cm−1 between the electronic ground and the first excited state, which is again in the same order as the two-orbital value. In order to draw a further comparison between the two-orbital model and the NEVPT2 calculations a scan along the mode of distortion for {[**Mo**]N}− was performed. A detailed description of how to perform such a scan is given elsewhere.<sup>5</sup> Relative energies of the four lowest electronic states as well as molecular g-tensors were calculated with the  $NEVPT2(7,8)$  method at each point of the scan (Figure 9). Note that this slightly reduced active space was chosen for the scan because it still provides a balanced description of the metal-ligand bonding but does not cause as many problems with convergence as the full 13 electron in 11 orbitals active space. As expected, the degeneracy of the four <sup>2</sup>E states is lifted even for the approximately  $C_{3V}$ -symmetric structure due to SOC and the imperfect trigonal symmetry of the complex geometry. The first excited Kramer's pair lies ~1000 cm<sup>-1</sup> above the lowest pair. Thus, one is close to true orbital degeneracy and hence linear response or perturbation theory would be invalid for the calculation of the g-tensor, as indicated by the very large deviations of the g-values from the free-electron value (in fact, linear response DFT yields g-values for the complete structure of  ${[\textbf{Mo}]}$ <sup>-</sup> yields values of  $g_1 = 1.99$ ,  $g_2 = 1.98$ ,  $g_3 = 1.78$  that are in strong disagreement with experiment).

The calculated g-values follow closely the trend predicted by the two-orbital model (Figure 7). The g-tensor remains axial at all points of the scan. Furthermore the three g-values evolve similarly to the  $m_l = \pm 2$  case in Figure 7. Thus the physical picture drawn for the gtensor above within the two-orbital model agrees with the quantum chemical calculations.

However, for the limiting cases of an undistorted and a fully distorted geometry, the NEVPT2(7,8) calculations numerically differ considerably from the two-orbital values. The  $g_{\parallel}$ -value at the C<sub>3V</sub> symmetric structure is calculated to be 0.35, which is significantly lower

than the value of 2 predicted by two-orbital model. Furthermore the  $g_{\perp}$  value is predicted to be 0.22, whereas it is predicted to be equal to zero by the two-orbital model. These findings have two causes: a) strong covalency present in the {**[Mo]**N}− complex and b) coupling of SOC to higher excited d-d states that are not included in the two-orbital model. The NEVPT2(7,8) calculations on  $\{[Mo]N\}^-$  at the distorted geometry predict g-values of  $g_1 =$ 1.91  $g_2$  = 1.90 and  $g_3$  = 1.39. While the first two g-values match the experimentally observed values almost perfectly, the third value is predicted significantly lower. Enlargement of the active space to 13 electrons in 11 orbitals leads to improved values of  $g_1$  $= 1.91$ ,  $g_2 = 1.91$  and  $g_3 = 1.48$ . However, the deviation of  $g_3$  from the experimental value is still not negligible. However, the results are considered to be satisfactory given the very steep dependence of the g-values of the precise energetic position of the first excited state (compare Figure 9). In the calculations this will depend on many details including basis set, higher order contributions of dynamic electron correlation, geometry, treatment of the environment, possible contributions from vibronic coupling, the precise treatment of relativistic effects among many others. All of these factors can change the calculated excitation energy by a few hundred wavenumbers. This is inconsequential for the g-tensor if the excitation energy is as large as several thousand wavenumbers but does show up prominently in the results if it is less than a thousand wavenumbers. From this perspective, the CASSCF(13,11)/NEVPT2 results are excellent.

It is known that CASSCF consistently underestimates covalency which in the present case acts to increase the g3 value. In the framework of the previously applied DDCI2 method dynamic correlation effects are taken into account for the computation of all three terms in equation (4).<sup>5,43,44</sup> As a result, a DDCI2(1,5) calculation of the model structure of {[**Mo**]N}− yields an increased g3 value of 1.55, while the other two values remain almost unaffected ( $g_1 = 1.92$ ,  $g_2 = 1.91$ ). Thus these results are internally consistent and are considered to be in good agreement with experiment. We thus believe that the electronic ground state description provided by the applied multireference methods and the physical picture behind the molecular g-values presented above is largely correct. Nevertheless it is interesting to note that the DDCI2(1,5) calculations yield values closer to the experiment than even the NEVPT2(13,11) calculations. We tentatively attribute this observation to the fact that the former uses the entire correlated wavefunctions and densities during the evaluation of SOC matrix elements, whereas the latter only employs spin-densities on the CASSCF(13,11) level together with NEVPT2(13,11) state energies.

The spin density distribution of the electronic ground state of {[**Mo**]N}− obtained from the DDCI2(1,5) wavefunction reveals that the spin is mainly located in the 4e manifold (Figure 10). However, its considerable tilt out of the equatorial plane reflects, as above, the distortion induced mixing of states with  ${}^{2}E_{b}$  and  ${}^{2}E_{a}$  parentage. The averaged metal d-character of the natural DDCI2 orbitals that correspond to the 4e orbital manifold is 81% which is in good agreement with 78% metal character obtained from the CASSCF(13,11) calculations.

These values strongly differ from the covalency parameter  $k = 0.25$  obtained from the twoorbital model. This highlights that the two-orbital model oversimplifies the electronic structure by absorbing multiple complex effects such as covalency and the aforementioned excited state-mixing into the single parameter k. As a result this parameter is artificially

reduced to an unphysical value of  $k = 0.25$ . The more realistic value of around 80% indicates that the bonding of the metal  $d_{x^2-y^2}/d_{xy}$  orbitals to the equatorial amido ligands is less covalent than the bonding between the  $d_{xz}/d_{vz}$  orbitals and the axial imido ligand in [**Mo**]NH.

#### **Quantum Chemical Computations of Hyperfine Interaction Parameters—**

Although it has to be stressed that orbital covalency is not a physical observable but a chemical concept to describe the nature of a chemical bond, hyperfine coupling constants and quadrupole interaction parameters provide a direct probe of the spin and charge density distributions. Thus, these values as calculated for the nitride and imido ligands of {[**Mo**]N}<sup>−</sup> and [**Mo**]NH within the linear response DFT framework are presented in Table 4, and discussed below in the context of the corresponding quantities as estimated by ENDOR spectroscopy. We have resorted here to DFT calculations since the CASSCF wavefunction, even the largest ones, are missing spin-polarization and hence provide unrealistic values for the important Fermi contact contribution to the hyperfine coupling. While, in principle, this can be remedied by higher-order dynamic correlation treatments such as NEVPT2, the NEVPT2 response density is not available and hence one would also only obtain CASSCF quality results. Except for the slight overestimation of covalency, the DFT spin-densities for the ground state are expected realistic since they provide a qualitatively similar description of the metal-ligand bonds as the more elaborate multireference calculations.

**14,15N ENDOR for [Mo]NH and {[Mo]N}−—**The molybdenum ions of {[**Mo**]N}−/ [**Mo**]NH have five nitrogen ligands in the first coordination sphere: the three amido nitrogen's in a plane perpendicular to the molecular axis plus the axial amino nitrogen, all associated with TAA, along with the axial nitride/imido ligand. However, with the aid of isotopic labeling the nitride and imide nitrogens can be identified.

**{[Mo]N}−:** Figure 11a shows the nitrogen Davies ENDOR spectra for {[**Mo**] <sup>14</sup>N}−, and 50% labeled  $\{[\text{Mo}]^{15}N\}^-$  collected at  $g_{\perp} = 1.91$ . As with the MoN<sub>2</sub> complex,<sup>5</sup> the dynamics of the JT effect lead to poorly-resolved 14/15N ENDOR spectra. Nevertheless, two distinct peaks are seen in the  ${[\text{Mo}]^{14}N}$ <sup>-</sup> spectrum at 5.3 and 8.5 MHz, riding on a broad background feature, and lose intensity in the spectrum from 50% labelled {[**Mo**] <sup>15</sup>N}−. Assignment of these as the  $v_{++}$  doublet (eq 2) from <sup>14</sup>N nitride split by the quadrupole interaction,  $|3P_{\perp}| = 3.2$  MHz, leads to a <sup>14</sup>N nitride hyperfine coupling,  $|A(^{14}N)_{\perp}| = 5.8$ MHz. This assignment correctly predicts the appearance of a single feature for the labeled  $\{[\text{Mo}]^{15}\text{N}\}^-$  at  $v_+ = 9.6$  MHz, based on the relation,  $|g_n|^{\frac{15}{N}}/g_n(^{14}\text{N})| = 1.4 = |A(^{15}\text{N})/g_n(^{14}\text{N})|$ A(<sup>14</sup>N). The absolute sign of the spin density on N,  $\rho(N) \propto A_{\perp}({}^{15}N)/g_n({}^{15}N) < 0$ , was determined by the Variable Mixing Time (VMT) PESTRE technique (Figure SI 3). A negative spin-population is consistent with the DFT calculations that predict −0.07 unpaired electrons on the nitrido ligand (Löwdin analysis). Although it is not possible to resolve features from the other coordinating nitrogens, we can estimate the  $^{14}N$  hyperfine couplings from the TAA ligands to be less than 6 MHz. This finding is consistent with the DFT calculations that predict hyperfine couplings for the amido nitrogens of 3–7 MHz and the amino nitrogen of <1 MHz.

The  $v_{+}$  peak from<sup>15</sup>N nitride of  $\{[\text{Mo}]^{14,15N}\}$ <sup>-</sup> in ENDOR spectra collected away from  $g_{\perp}$ , quickly broadens and moves towards low frequency and becomes lost (Figure SI 1) near  $g_{\parallel}$ , with the <sup>14</sup>N peaks showing a similar trend, behavior demonstrating that  $|A_{\parallel}(N)| \ll |A_{\perp}(N)|$ . The quantum chemical calculations predict a tensor  $(A<sub>nitride</sub>(1<sup>4</sup>N)= [-8 -2 1])$  MHz with small isotropic negative coupling ( $A_{iso} \sim -3$  MHz), in agreement with the ENDOR and VMT-PESTRE measurements. As shown in Figure SI 2, simulation of the 2D pattern of spectra collected at multiple fields with simulations that employ the calculated tensor are indeed in satisfactory agreement with the experimental data.

It is also possible to estimate the quadrupole interaction parameter for the  $^{14}N$  nitride,  $\vert$  $e^{2}qQ/h$  from the spectrum of Figure 11a. In the near-trigonal symmetry of the complex, the quadrupole interaction is expected to be effectively axial, with the unique value,  $P_{\parallel}$ , along  $g_{\parallel}$ . The quadrupole tensor is traceless,  $P_{\parallel}$  + 2P<sub>⊥</sub> = 0, and thus, the measurement of  $|P_{\perp}|$  yields the interaction parameter,  $|e^2qQ/h| = 4|P_\perp| = 4.2 \text{ MHz}$ . This parameter reflects the difference between the charge density in the nitride  $2p\sigma$  orbital along the Mo-N bond (N<sub>3</sub>) and the average of the densities in the two orthogonal  $2p\pi$  orbitals, and the experimental and calculated value, e <sup>2</sup>qQ/h = −3.47 MHz, are in excellent agreement (Table 4). In SI we present a simplified, heuristic discussion of these bonding interactions through use of the Townes-Dailey model, in which the quadrupole parameter for Mo-bound nitride is simply proportional to the difference between the charge density in the nitride  $2p\sigma$  orbital along the Mo-N bond  $(N_3)$  and the average of the densities in the two orthogonal  $2p\pi$  orbitals.

[Mo]NH: Although the ENDOR spectra of [Mo]<sup>14</sup>NH and 50% labelled [Mo]<sup>15</sup>NH measured at  $g_2 = 1.96$  are rich in peaks, there is no observable difference between them that can be assigned to the imido nitrogen, Figure 11b. The ENDOR spectra of [**Mo**] <sup>14</sup>NH, collected away from  $g_2$  quickly become featureless (Figure SI 4), which compounds the difficulty in assigning an imido nitrogen signal. The quantum chemical calculations predict a highly anisotropic hyperfine tensor  $(A = [30, 0, 0]$  with  $A_{iso} = 10$  MHz) for the imido nitrogen. Simulations of the ENDOR spectra predicted with this tensor are spread over the frequency scale (see Figure SI 6) without sharp features. This, along with the poor resolution of the spectra, may help to rationalize the failure to observe the nitride signals.

Although the background  $14N$  ENDOR signals from other coordinating nitrogens are unresolved, we can estimate that  $14N$  hyperfine couplings from the TAA ligand are less than 6 MHz. This is in agreement with the calculations, which predict amido nitrogen HFCs along the principle g-tensor axes of 3–7 MHz, with an even smaller coupling to the amino nitrogen of around 2 MHz. Interestingly, the <sup>14</sup>N ENDOR spectra of MoNH show additional peaks at frequencies above 10 MHz in the range of 10–16 MHz which are not observed in MoN−. For example, the peak at around 11 MHz is a double quantum peak of nitrogen with hyperfine coupling of A∼3 MHz (Figure SI 5). This could well be from one of the in-plane amido nitrogens.

#### **1,2H and 95,97Mo ENDOR for [Mo]NH and {[Mo]N}<sup>−</sup>**

**1,2H ENDOR:** Figure 12 shows 1H DAVIES ENDOR spectra for the two complexes recorded at the respective maxima  $(g<sub>⊥</sub>)$  of their EPR spectra. The proton ENDOR features

for {[**Mo**]N}− are centered at the 1H Larmor frequency, and as expected, exhibit signals from protons of the HIPT ligands, with hyperfine couplings of less than 5 MHz.50 In addition, a weak, broad peak centered at −12 MHz relative to the proton Larmor frequency is observed, which can be assigned as the  $v^{+} \sim 44$  MHz branch of the m<sub>I</sub> =  $\pm$  1/2 doublet for the  $I = 5/2$  <sup>95,97</sup>Mo ion, as discussed below.

The weakly-coupled 1H features from the HIPT ligand are also observed for [**Mo**]NH, as expected. The spectrum for  $[Mo]NH$  also shows intense <sup>1</sup>H signals with significantly stronger coupling; the broad  $v^+ \sim 7$ –17 MHz feature of a strongly-coupled <sup>1</sup>H, plus its  $v^$ partner, which is distorted in part because of overlap with a signal from  $95,97$ Mo. The assignment of this broad  ${}^{1}H$  feature to the imido proton is confirmed by the  ${}^{1}H$  spectrum of the <sup>2</sup>H labeled sample, MoN <sup>2</sup>H: H/D exchange abolishes the broad <sup>1</sup>H  $v_+ / v^-$  doublet but leaves both the 1H features from HIPT and the weak, broad 95,97Mo peak centered at −10 MHz relative to the proton Larmor frequency (fig 12).

The 2D pattern of 1H ENDOR spectra collected across the full EPR envelope for [**Mo**]NH (Figure SI 7) show two resolved broad features at fields between  $g_1(1.98)$  and  $g_2(1.96)$ , suggesting two major but poorly defined NH conformers have been trapped, with couplings at  $g_1$  of  $\sim$  22 and 30 MHz. These features merge and the coupling decreases as the field is increased towards  $g_3 = 1.86$ , where  $A_3 \sim 18(6)$  MHz. Precise hyperfine tensors cannot be obtained from the broad and poorly-resolved features of the overlapping 1H responses from the two  $N^1$ H conformers, but we can estimate that each has an isotropic coupling of roughly,  $|A_{\text{iso}}| \sim 20$  MHz.

**95,97Mo ENDOR:** The <sup>95,97</sup>Mo ENDOR response (\*) seen at g⊥ for {[**Mo**]N}<sup>−</sup> (Figure 12) corresponds to a hyperfine coupling of  $|A^{(95,97)}Mo|_1 \approx 80$  MHz. Although the possibility that the relatively narrow width of this feature is indicative of a relatively small quadrupole interaction, it is by far more likely that the interaction is large and this signal corresponds to the  $m_1 = +1/2 \leftrightarrow -1/2$  transition, and that additional branches associated with the quadrupole-split  $m_I = \pm 3/2$ , 5/2 features are not observable. In spectra collected across the EPR envelope (Fig SI 12), the <sup>95,97</sup>Mo signal for  $\{[\text{Mo}]\}$ <sup>-</sup> is quickly lost away from g⊥, likely because of strong anisotropy in the couplings. The calculations in fact show a strong anisotropy. The calculated hyperfine tensor is essentially axial, with  $A(^{95,97}Mo_{\perp} \approx 77$ MHz, in excellent agreement with experiment, and  $A(^{95,97}Mo)_{\parallel} = 167 \text{ MHz}.$ 

The <sup>95,97</sup>Mo feature for { $[Mo]NH$ } in the g $\perp$  region also corresponds to a hyperfine coupling  $A(^{95,97}Mo)$ <sub>⊥</sub> ≈ 80 MHz (Figure 12), essentially the same as that for  $\{[Mo]N\}^-$ . For {[Mo]NH} the<sup>95,97</sup>Mo signal is visible, and shifts little over most of the EPR envelope, between  $g_1 = 1.98$  and  $g \sim 1.90 \rightarrow g_3$ , (Figure SI 7), suggesting that the <sup>95,97</sup>Mo coupling is largely isotropic,  $A(^{95,97}Mo)_{iso} \approx 80 \text{ MHz}$ . The calculations yield an isotropic coupling that matches this quite well,  $A(^{95,97}Mo)_{iso} = 77 MHz$ , but shows an appreciable anisotropy.

#### **Discussion**

In this work we have presented a combined spectroscopic and theoretical study of the catalytically active **[Mo]**NH complex and its derivative **[Mo]**N−. On the basis of our findings

we were able to establish a coherent picture of the complex electronic structure and magnetic properties of both these Mo(V) compounds, thus completing an in-depth characterization of the four states of Figure 1. The tripodal trisamidoamine ligand together with the axial amido- and imido- moieties creates an idealized  $C_{3V}$ –symmetric ligand field around the Mo(V) centers. In the case of **[Mo]**NH, our correlated multireference quantum chemical calculations predict that the imposed ligand field results in an  ${}^{2}E$  electronic ground state that features a half-filled e(xz,yz) shell. This prediction is in agreement with previous findings made for analogous  $Mo(III)$  and similar Fe complexes.<sup>13,14</sup>

As observed previously for  $[Mo]NH_3$  and  $[Mo]N_2$ , the half-occupation of the orbitally degenerate e(xz,yz) shell and the presence of unquenched spin-orbit coupling effects leads to a Pseudo-Jahn-Teller behavior.<sup>5</sup> The competing effects of spin-orbit and Jahn-Teller coupling result in a delicate equilibrium that is effectively probed by our measurements of the molecular g-values. In-state spin-orbit coupling forces the g-values, in particular the component along the molecular z-axis, away from free electron value, whereas the Jahn-Teller distortion has the opposite effect by lifting this (near) degeneracy. Hence, the experimentally observed molecular g-values give insight into the relative strength of both effects. Our analysis of the experimental data in the framework of a 2-orbital model even allows for an assignment of a numerical value to the relative strength. For **[Mo]**NH we find a strong Jahn-Teller splitting states resulting in two well separated states with  ${}^{2}E$  parentage. Moreover, the analysis indicates a strong mixing of metal and ligand orbitals for the  $e(xz,yz)$ manifold.

Our CASSSCF/NEVPT2 calculations of the molecular g-tensor for **[Mo]**NH is in excellent agreement with the observed experimental values thus confirming the overall picture of the electronic structure, while give deep insights into details of that structure. Interestingly, almost the same numerical agreement between experiment and theory is achieved with simple DFT calculations. We believe that this observation is due to the clear separation of the electronic ground state from the first excited state. Generally, only correlated multireference calculations provide the required flexibility and accuracy to reliably predict the electronic structure of such complicated cases. Another consequence of the strong Jahn-Teller coupling is the distortion-induced mixing of states with  ${}^{2}E(xy, x^{2}-y^{2})$  parentage to the electronic ground state. This mixing, which by construction is not described in our 2-orbital model, is mostly responsible for the rhombicity of the molecular g-tensor. Furthermore, as the mixing shifts the HOMO out of the sterically overloaded x-y plane, it may facilitate the attack of the imido nitrogen by an electrophile such as a proton. A similar argument has been made previously for tripodal Fe compounds.<sup>18</sup>

Our calculations predict that the {**[Mo]**N−} precursor to **[Mo]**NH in the catalytic cycle has a qualitatively different electronic ground state. For {**[Mo]**N−}, the electronic ground state is still a <sup>2</sup>E-state but with a half-filled  $e(xy, x^2-y^2)$  shell. This change is induced by the extraordinarily strong ligand field along the molecular z-axis created by the amido ligand. Despite the qualitative differences to **[Mo]**NH, we again find a Pseudo-Jahn-Teller behavior with an equilibrium of spin-orbit and Jahn-Teller coupling for {**[Mo]**N−}, although with slightly altered physical parameters. Analysis of the experimentally observed molecular gvalues within the 2-orbital model indicates that the equilibrium is tilted towards the spin-

orbit coupling compared to **[Mo]**NH. This finding is intelligible considering the fact that the  $e(xy, x^2-y^2)$  shell corresponds to the M =  $\pm 2$  manifold of the Mo d-orbitals whereas the e(xz,yz) shell corresponds to the  $M = \pm 1$  manifold. Our CASSCF/NEVPT2 calculations of the molecular g-values again achieve good agreement with the experimentally observed values thus validating our in-depth electronic structure description.

It is important to note that the change of character of the electronic ground state is likely to be accompanied by a change of reactivity. For example, the shift of the HOMO out of the xy plane into the z-direction may lead to a lowering of the barrier for an electrophilic attack at the nitride ligand.

The formal two-orbital model indicates that the parameter  $r$ , the ratio of  $JT$  coupling to SOC (**eq 9**), is ~ two-fold larger for **[Mo]**NH than for {**[Mo]**N−}. However, within this model, according to **eq 9**, the JT coupling is in fact ~ four-fold greater for **[Mo]**NH, because in the denominator the factor k for the SOC of **[Mo]**NH is twice that for {**[Mo]**N−}. The large JT coupling reflects the non-linearity of the Mo-N-H moiety, as discussed above. Comparing the parameters, r, for the three-fold d<sup>1</sup>{[Mo(V)]N<sup>-</sup>} with those of the three-fold d<sup>3</sup> **Mo(III)-L, L= N<sub>2</sub>, CO, and NH<sub>3</sub><sup>5</sup> once one takes in into account the differences in k, one** finds that the JT coupling is essentially the same for the two configurations, despite the difference in ground state, 4E for the former versus 3E for the latter (Figure 4). The latter is significant in terms of the SOC interactions inside the given E-term. For the 4e-set of orbitals, which is based on the Mo  $d_{xy}$  and  $d_{x2-y2}$  orbitals, the angular momentum matrix element along the z-axis is +/−2, whereas for the 3e set, which is comprised of the Mo based  $d_{xz}$  and  $d_{yz}$  orbitals, the corresponding matrix element is +/−1. Hence, effectively, the SOC interaction in the 4E term is twice as large as in the 3E term. This general trend is modulated by metal-ligand covalency. Thus, for {**[Mo(V)]**N−} vibronic coupling is induced by partial occupation of orbitals with mostly  $d_{xy}/d_{x^2-y^2}$  character that are involved in the  $\sigma$ -type bonding of the Mo center to the equatorial amide ligands in {**[Mo(V)]**N−}. In contrast, the origin of vibronic coupling in the three  $d^3$  complexes and [Mo]NH is the partial occupation of orbitals with mostly  $d_{xx}/d_{yz}$  character that facilitate  $\pi$ -bonding of the Mo center to the axial ligands. Moreover, one has to keep in mind that for {**[Mo(V)]**N−} SOC acts to lower the g<sub>II</sub> value whereas it leads to an increase of g<sub>II</sub> for **Mo(III)-L**, **L**= N<sub>2</sub>, CO, and NH<sub>3</sub>.<sup>5</sup>

In addition to the molecular g-tensor we have measured the  $^{14,15}$ N and  $^{95,97}$ Mo ENDOR of **[Mo]**N and 1,2H, 14,15N and 95,97Mo ENDOR of **[Mo]**NH. The 14,15N hyperfine coupling measured by ENDOR for **[Mo]**N is small, in keeping with the orthogonality of both the σ and  $\pi$  orbitals on N to the 4e spin-bearing orbitals on Mo. The experimental and calculated  $14N$  hyperfine couplings are in excellent agreement. The negative sign of  $a<sub>iso</sub>$ for 14N, as both calculated and determined by experiment, corresponds to delocalization through spin polarization of the Mo-N bonding orbitals. The measured quadrupole coupling is in excellent agreement with the calculated value, and indicates  $\sigma$  donation from N to Mo is much more significant that  $\pi$  back-donation to N. It may be noted that the bonding in an analogous trigonal  $d^3$  Fe(V)≡N complex appears to be quite different. Whereas the Mobound nitride has a distinct difference in the occupancies of nitrogen  $2p\sigma$  versus  $2p\pi$ orbitral, analyses of the 14N quadrupole and hyperfine couplings of the Fe-nitride ligand

support a surprising picture in which the  $Fe(V)$  ion binds an essentially spherically symmetric nitride trianion.<sup>18</sup> The <sup>14,15</sup>N hyperfine coupling for  $[Mo]$ <sup>14,15</sup>NH could not be resolved, and calculations suggest that this results from a highly anisotropic interaction tensor. However, the  $^{1,2}$ H signals were readily detected and interpretable with an anisotropic coupling tensor with substantial isotropic coupling, aiso( ${}^{1}$ H) ~ 20 MHz, with satisfactory agreement between computation and experiment.

Finally, we can use the present results to comment on the alternative pathways for [e−/H+] addition during the reduction of **[Mo(VI)]**N to **[Mo(V)]**NH during catalysis, Figure 1. The ability to trap and crystallographically characterize  $\{[\text{Mo}(V\text{I})|NH]\}^+$  clearly demonstrates that the 'proton-first' pathway is accessible during catalytic reduction of  $N_2$ . However, the present characterization by experiment and theory of {[**Mo(V)**]N}−, in the context of earlier calculations on the reaction mechanism, nonetheless suggest that the 'electron-first' pathway is feasible. $11$ 

#### **Summary**

Our analysis shows that Jahn-Teller and spin orbit coupling effects have a significant impact on the electronic structures of **[Mo]**NH and **[Mo]**N− and are absolutely key for understanding the physics and chemistry of these compounds. In particular the magnetic properties of these compounds are governed by the delicate balance of the two effects. Furthermore, together with our previous work on **[Mo]** complexes we have established the combination of EPR spectroscopy and correlated multirefence quantum chemical calculations as a valuable tool to probe and analyze the electronic structure of compounds with near-degenerate ground states.<sup>5 14,15</sup>N and <sup>1,2</sup>H hyperfine couplings from the axial –N and −NH ligands provide stringent tests of the validity of quantum calculations, and in fact calculation and experiment are in excellent agreement. The  $95,97$ Mo hyperfine couplings should provide a useful benchmark in considerations of the state of Mo in synthetic and biological multimetallic clusters. This becomes particularly relevant for the case of the FeMo-cofactor catalytic cluster of nitrogenase, where the oxidation state Mo(III) has recently been suggested for the Mo center on the basis of spectroscopic experiments and quantum chemical calculations.<sup>51</sup>

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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#### **Synopsis**

The molybdenum trisamidoamine (TAA) complex [**Mo**] (=(3,5-(2,4,6-i-Pr3C6H2)2C6H3NCH2CH2N)Mo) carries out catalytic reduction of N2 to ammonia by protons and electrons at room temperature. A key intermediate in the proposed [**Mo**] nitrogen reduction cycle is nitrido-Mo(VI), [**Mo(VI)**]N: the addition of [e−/H+] to [**Mo(VI)**]N generates [**Mo(V)**]NH In combination with previous X-ray studies, EPR/ ENDOR and multli-reference based computational studies of {[**Mo(V)**]N}− and [**Mo(V)**]NH create the rare situation where each one of the four possible states of an [e−/H+] delivery has been characterized. Jahn-Teller and spin orbit coupling effects have a significant impact on the electronic structures of **[Mo]**NH and **[Mo]**N− and are absolutely key for understanding the physics and chemistry of these compounds



#### **Figure 1.**

The three possible pathways for addition of [e−/H+] to [**Mo**]N As shown, and discussed in text, two states have been structurally characterized; two are characterized here by paramagnetic resonance and computational methods.



#### **Figure 2.**

Model structure of **[Mo]**N−. The HIPT groups have been replaced by smaller ethyl groups, which retain an electron donating effect but are computationally less demanding.





#### **Figure 3.**

Active orbitals of {[**Mo**]N}− (left) and [**Mo**]NH (right) in CASSCF(7,8) and NEVPT2(7,8) calculations. Both sets of orbitals are obtained as natural orbitals of converged CASSCF(7,8) calculations. Note, in the case of  $[Mo]NH$  the  $d_{z^2}$  and its bonding counterpart as well as the  $d_{xy}$  orbital are not contained in the active space. The enumeration of the orbitals refers to the decomposition into contributions from atomic orbitals given in Table 1 and Table 2.



#### **Figure 4.**

Schematic MO diagram of  $[Mo]NH$  (a) and  $[Mo]N^-(b)$ . In both cases the  $4a_1(z^2)$  orbital is strongly destabilized by the axial ligand. The energetic order of the  $3e(xz,yz)$  and  $4e(x^2-z)$  $y^2$ ,xy) orbitals changes with the loss of the proton due to the different ligand field strengths of the imido and the nitrido ligands (see text).



#### **Figure 5.**

Modes of distortion for [Mo]NH. In the equatorial plane,  $N_2$  and  $N_3$  bend towards  $N_1$ resulting in reduced angles of ∠ ( $N_1M_0N_3$ ) = 109° and ∠ ( $N_1M_0N_2$ ) = 112°. Consequently, the angle ∠ ( $N_2M_3$ ) = 123° is significantly larger. Note, the sum of all three angles does not equal 360° as the angles were determined from the three-dimensional structure rather than a two-dimensional projection. As indicated in the left and middle panel the axial nitrogen moves towards  $N_3$ . Interestingly, the adjacent hydrogen slightly bends towards the gap between N1 and N3. For {[**Mo**]N}− an analogous combination of equatorial bending and transversal motion is observed.



#### **Figure 6.**

Field swept ESE-EPR for {[**Mo**]N}− and [**Mo**]NH. The (\*) represents the epr signal at g ~ 2 of matrix free radicals generated during cryoreduction. For clarity, the epr signal of the radical is cut off. The unique g values  $(g_{\parallel}, g_{\perp})$  are shown with arrows. Conditions, two pulse echo detected EPR: microwave frequency,  $34.881-34.965$  GHz;  $\pi = 80$  ns;  $\tau = 700$  ns; repetition time, 100 ms.





#### **Figure 7.**

Evolution of the molecular g-values for a singly occupied orbital with  $m_l = \pm 1$ , as appropriate for [Mo]NH and  $m_1 = \pm 2$ , as for {[Mo]N}<sup>-</sup>, with increasing PJT splitting within the two-orbital model and the ionic limit of  $k = 1$ . For convenience, an ionic SOC constant of  $\lambda_{\text{SOC}}$  = 800 cm<sup>-1</sup> was used for both. Within the two-orbital model, the g-tensor is predicted to be axial at all points.



#### **Figure 8.**

Electron Spin density of the electronic ground state of [**Mo**]NH obtained from a DDCI(1,5) calculation at an isovalue of 0.003 electron spins.



#### **Figure 9.**

Evolution of the nonrelativistic NEVPT2(7,8) energy (left panel) of the lowest two Kramer's pairs, the difference of the first energy of the lowest Kramer's pair (calculated on the NEVPT2(7,8) level) after inclusion of spin-orbit coupling (middle panel) and the molecular g-values (right panel) for {[**Mo**]N}− with increasing distortion of the molecular geometry from  $C_{3V}$  symmetry. More precisely, the quantity  $\rho$  on the x-axis of all three panels is the displacement along the Jahn-Teller active interaction coordinate (see above). Note that the lines for  $g_2$  and  $g_3$  are virtually indistinguishable at all points of the scan, which corresponds to an axial g-tensor as predicted within the 2-orbital model (Fig 5)



#### **Figure 10.**

Electron Spin density of the electronic ground state of {[**Mo**]N}− obtained from a DDCI2(1,5) calculation at an isovalue of 0.003 electron spins. The tilt out of the equatorial plane reflects the mixing of states with  ${}^{2}E_{b}$  and  ${}^{2}E_{a}$  parentage.



#### **Figure 11.**

**(a)** Davies 14,15N ENDOR of  $\{[\text{Mo}]^{14}N\}^-$ , and 50%<sup>15</sup>N labeled  $\{[\text{Mo}]^{14,15}N\}^-$  at  $g_{\perp} =$ 1.91. The quadruple splitting for the nitride nitrogen is shown as 3P; nuclear Larmor frequencies,  $v_{14N/15N} = 3.9/5.5 \text{ MHz}$ , **(b)** Davies <sup>14,15</sup>N ENDOR of  $[Mo]$ <sup>14</sup>NH, and 50%<sup>15</sup>N labeled [Mo]<sup>14,15</sup>NH at  $g_2 = 1.96$ . Conditions: microwave frequency, 34.881– 34.965 GHz;  $\pi = 200$  ns;  $\tau = 700$  ns;  $t_{\text{rf}} = 30$  µs; repetition time, 50 ms;



#### **Figure 12.**

Davies <sup>1</sup>H ENDOR of  $\{[\text{Mo}]\text{N}\}^-$ ,  $[\text{Mo}]\text{NH}$  (each black), and  $[\text{Mo}]\text{ND}$  (red) at  $g_{\perp} = 1.91$ , 1.96 respectively. For [**Mo**]NH, the brace shows the broad distribution of the proton coupling. The broad feature (\*) observed at −12 MHz is contribution from 95,97Mo as explained in the text. Conditions: microwave frequency, 34.881–34.965 GHz;  $\pi$  = 120 ns;  $\tau$  $= 700$  ns; t<sub>rf</sub> = 60 µs; repetition time, 50 ms; RF frequency randomly hopped.



**Chart 1.**  The [**Mo**]NH complex

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## **Table 1**

Important atomic orbital contributions to the active orbitals in CASSCF(7,8) and NEVPT2(7,8) calculations on the ethyl model of {[Mo]N]<sup>-</sup>. All Important atomic orbital contributions to the active orbitals in CASSCF(7,8) and NEVPT2(7,8) calculations on the ethyl model of {[**Mo**]N} contribution are given in %. contribution are given in %.



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# **Table 2**

Important atomic orbital contributions in to the active orbitals in CASSCF(7,8) and NEVPT2(7,8) calculations on the ethyl model of [**Mo**]NH. All Important atomic orbital contributions in to the active orbitals in CASSCF(7,8) and NEVPT2(7,8) calculations on the ethyl model of [Mo]NH. All contribution are given in  $\%$  . contribution are given in %.



#### **Table 3**

Excitation energies and dominant configurations of the d-d multipltes for **[Mo]**N− and **[Mo]**NH from NEVPT2(13,11) calculations as described in the text. An excitation energy of 0 cm−1 signifies the electronic ground state.



a)These excitation energies have been determined by NEVPT2(1,5) calculations because of interfering CT states in the NEVPT(13,11) calculations.

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## **Table 4**

Calculated nuclear hyperfine parameters that were obtained using the B3LYP functional together with the TZVP basis set and the ZORA Hamiltonian on Calculated nuclear hyperfine parameters that were obtained using the B3LYP functional together with the TZVP basis set and the ZORA Hamiltonian on the full [Mo]X (X=N<sup>3-</sup> and NH<sup>2-</sup>) model. The given Euler angles refer to the principle axis system of the molecular g-tensor. the full [Mo]X (X=N<sup>3−</sup> and NH<sup>2−</sup>) model. The given Euler angles refer to the principle axis system of the molecular g-tensor.



#### **Table 5**

Experimentally observed and calculated bond angles for [**Mo**]N and [**Mo**]NH.

