

High-Spin Iron(VI), Low-Spin Ruthenium(VI), and Magnetically Bistable Osmium(VI) in Molecular Group 8 Nitrido Trifluorides NMF₃

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Abstract: Pseudo-tetrahedral nitrido trifluorides N≡MF₃ (M = Fe, Ru, Os) and square pyramidal nitrido tetrafluorides N≡MF₄ (M=Ru, Os) were formed by free-metal-atom reactions with NF₃ and subsequently isolated in solid neon at 5 K. Their IR spectra were recorded and analyzed aided by quantumchemical calculations. For a d² electron configuration of the $N \equiv MF_3$ compounds in C_{3v} symmetry, Hund's rule predict a high-spin ${}^{3}\text{A}_{2}$ ground state with two parallel spin electrons and two degenerate metal $d(\delta)$ -orbitals. The corresponding high-spin ³A₂ ground state was, however, only found for N≡FeF₃, the first experimentally verified neutral nitrido Fe^{VI} species. The valence-isoelectronic N≡RuF₃ and N≡OsF₃ adopt different angular distorted singlet structures. For N≡RuF₃, the triplet ³A₂ state is only 5 kJ mol⁻¹ higher in energy than the singlet ¹A' ground state, and the magnetically bistable molecular N≡OsF₃ with two distorted near degenerate ¹A′ and ³A" electronic states were experimentally detected at 5 K in solid neon.

Introduction

The group 8 transition metals have eight electrons in their valence shell, but in addition to the well-known strong oxidizers RuO₄ and OsO₄, only Os has a variety of different complexes in oxidation state VIII.[1] While the oxidation state VI is abundant for ruthenium and osmium, the complex anion [FeO₄]²⁻ was the only known Fe^{VI} compound for a long time. [2] In 2007 the neutral, dioxo Fe^{VI} peroxide $O_2Fe(\eta^2-O_2)$ was reported to be formed from molecular FeO₂ and O₂ under cryogenic conditions.[3] Tetrahedral Fe^{VIII}O₄ was shown to be metastable with respect to $O_2Fe^{VI}(\eta^2-O_2)$ in the gas phase, [4] and the oxidation state VII is so far the highest oxidation state of iron observed experimentally for the tetrahedral tetroxide anion FeO₄-.^[5] In addition to oxygen, nitrogen ligands are also able to stabilize high oxidation states of iron. Such terminal iron-nitrido complexes have already been the subject of several up-to-date reviews.^[6] We restrict ourselves to some representative examples such as the square-pyramidal [(TPP)Fe V N] (TPP $^{2-}$ = tetraphenylporphyrinate dianion), characterized by spectroscopy, [7] the tetragonal nitrido FeVI dication [(Me3cyac) $([Me_3cyac]^- = N,N,N-tri-methyl-1,4,8,11-tetraazacyclote$ tra-decane-1-acetate), confirmed by Mössbauer and X-ray spectroscopy, [8] the pseudo-tetrahedral [(PhB(PCH₂PiPr₂)₃)Fe^{IV}N] $(PhB(PCH_2PiPr_2)_3 = tris(diisopropylphosphinophenyl)borane),^{[9]}$ and, very recently, the crystal structure of a thermally stable four-coordinate Fe^{VI} bis(imido) cation, [(H₂B(MesIm)₂)Fe(= $NMes_{2}$ ⁺ ($[H_{2}B[MesIm]_{2}]^{-}$ = dihydrobis-[1-(2,4,6-trimethylphenyl)]imidazol-2-ylidene]borato).[10]

Nitrido iron complexes play an important role in a number of chemical and biological processes, for example in the catalytic cycle of cytochrome P450,[11] in the FeMo cofactor of the nitrogenase enzyme^[6a] and in the Haber–Bosch process.^[12] In analogy to the active iron surface nitride in the Haber-Bosch process, ammonia synthesis has also successfully achieved under mild conditions using the ruthenium pincer nitrido complex [(PNP)RuN] $(PNP^- = [N(CH_2CH_2P^tBu_2)_2)]^-).^{[13]}$ Quite recently osmium(VI) nitrides have emerged as a new class of potential anticancer and antitumor agents.[14] Examples include $[(bipy)Cl_3Os^{VI}N]^{[15]}$ $(bipy=2,2'-bipyridine)^{[16]}$ and $[(sap)(py)]^{[16]}$ ClOs^{VI}N] (sap = deprotonated *N*-salicylidene-2-aminophenol).^[17] The wide field of possible applications of group 8 nitrido complexes underline the importance of a deeper understanding of the properties of this class of compounds. Especially the nitrido metal-ligand multiple bond and the valency of the metal are key factors for the reactivity and structure of these

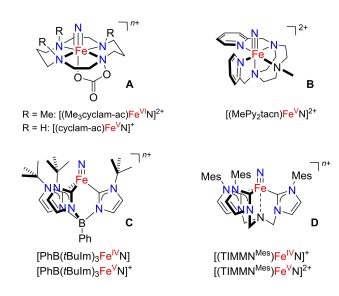
In particular, there has been a tremendous progress in the synthesis and the chemistry of molecular Fe^{IV} and Fe^{V} nitrido compounds in the recent years that have been described in detail in several review articles. [6] They are supported by sterically encumbered macrocyclic or chelating ligands involving nitrogen or N-heterocyclic carbene donors based on, for example porphyrin or nitrogen- and boron-anchored tri- and tetrapodal chelates to protect the reactive Fe=N moiety (see Scheme 1 for representative examples). The most common route to these nitrido compounds is the photolysis of an iron azido precursor and concomitant N₂ evolution, whereby the

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Scheme 1. Representative examples of ligand-supported tetragonal ($\mathbf{A}^{[8,22]}$ and $\mathbf{B}^{\text{[19d]}}$) and trigonal ($\mathbf{C}^{\text{[19a]}}$ and $\mathbf{D}^{\text{[19b]}}$) coordinated high-valent iron nitrido complexes.

one-electron oxidation of the Fe^{IV} nitrido complexes often represents an alternative route to Fe^V nitrido complexes. [6a,b]

The reactivity of these high-valent nitrido iron compounds in chemical transformations have been thoroughly explored, [6,18] their structures, and their electronic properties have been investigated in detail using a variety of experimental and quantum mechanical methods.[19] While these studies contributed greatly to the understanding of the iron nitride bonding motif, our knowledge about the behavior, the nature, and bond-strengths of the Fe≡N triple bond in high valent iron compounds upon iron oxidization is, however, still very limited and contradictory. Two questions arise here: Is there a nitrido wall[20] from which the nitrido ligand gives up its innocent behavior,[21] and does the Fe=N bond become stronger and stronger through oxidation of the iron center?

It should be emphasized that the known iron nitrido species can be divided into trigonal (pseudo-tetrahedral) and tetragonal (pseudo-octahedral) complexes (Scheme 1), since these two groups show different ligand field splitting of the Fe(3d) orbitals. [9,23] In a trigonal C_{3v} ligand field there are two purely Fe \equiv N nonbonding *e*-type orbitals (d_{xy,x^2-y^2}) , which allow the accommodation of up to four electrons energetically below the antibonding Fe=N orbitals. [6a,c,9,23] This results in a relatively strong Fe≡N triple bonds, for example, low spin Fe^{IV} derivatives, for which very short experimental Fe-N distances (Table S1 in the Supporting Information) and Fe-N stretching vibrations at 1008–1034 cm⁻¹ were found. [9,24]

Conversely, in the tetragonal C_{4v} ligand field there is only one purely nonbonding (d_{xy}) orbital with respect to the Fe \equiv N bond energetically below the π^* -antibonding $(d_{xz,yz})$ MOs. [6a,c] A d-electron count larger than two results here in the occupation of $\pi^*(Fe\equiv N)$ orbitals, and, accordingly, Fe^{IV} (d⁴) and Fe^{V} (d³) nitrido complexes in tetragonal symmetry are generally thermally less stable and more reactive. [6b,18a,c] Note that the d³ ground-state electron configuration of Fe^V nitrido complexes is subject to a Jahn-Teller distortion. [19b,c] To overcome the thermal instability and high reactivity of such tetragonal Fe^V nitride complexes their Fe=N distances and stretching frequencies were obtained by a variety of spectroscopic methods either at cryogenic temperatures or at the gas phase (for representative examples, see Table S1). As expected, the experimental Fe-N distances for the two tetragonal complexes [Fe^{v-} $(N)(MePy_2tacn)]^{2+}$ (Scheme 1, 3d⁴ configuration, Fe–N: 164(1) pm)^[19d] and [Fe^V(N) (cyclam-ac)]⁺ (Scheme 1, cyclam-ac= Fe-N: 1,4,8,11-tetraazacyclotetradecane-1-acetato, pm), [22] estimated from extended X-ray absorption fine structure (EXAFS) analysis, were found to be longer than the Fe-N distance of the analogous Fe^{VI} dication [Fe^{VI}(N)(Me₃cyclam-ac)]²⁺ (Scheme 1, 157(2) pm) with a singlet 3d² configuration.^[8]

In contrast, the formal Fe=N bond order in trigonal Fenitrido complexes does not change by increasing the iron oxidation state from singlet Fe^{IV} to triplet Fe^{VI}, making predictions about the bond lengths less intuitive as other factors such as the geometry and the nature of the ligands come to the fore. X-ray structure analysis of the $Fe^{IV}N/Fe^VN$ derivatives of the two redox pairs [PhB(tBulm)₃FeN]^{0/+} (Scheme 1) $^{[19a,24b]}$ and $[(TIMMN^{MES})FeN]^{+/2+}$ (Scheme 1) $^{[19b]}$ show different trends. While the Fe-N length decreases slightly from 151.2(1) pm to 150.6(2) pm for the former, it increases from 151.3(3) pm to 152.9(1) pm for the latter. The different trend in these Fe≡N distances during oxidation of Fe^{IV} to Fe^V was attributed to a possibly stronger interaction between the ligand N anchor with the more electrophilic Fe^V center in [Fe-(N)(TIMMN^{MES})]²⁺ (Scheme 1).^[19b] On the other hand, also coordinated solvent molecules can make it difficult to compare the Fe≡N distances of different complexes, since this leads to shortened experimental Fe \equiv N distances. [19e]

In this work, we describe the preparation of the molecular, neutral nitrido trifluorides $NM^{VI}F_3$ of the group 8 metals M = Fe, Ru, Os from IR laser ablated metal atoms and gaseous NF3 and their IR-spectroscopic characterization under cryogenic conditions in a noble gas matrix. These trigonal nitrido trifluorides bear genuine M

N triple bonds, unsupported by sterically encumbered electron donor substituents with the innocent fluoride ligand. The M≡N stretching vibration of theses derivatives is energetically sufficiently isolated from other fundamentals. Hence, it is considered to be a reliable experimental signature for M–N bond strength and M–N bond length in these nitrido complexes. This analysis overcomes the difficulties described above and also has the advantage that the experimental results can be supported and analyzed by reliable and accurate quantum mechanical calculations of these molecular, neutral compounds. Furthermore, this analysis enables a direct comparison of experimental M≡N stretching frequencies of M = Fe^{VI} and its heavier group 8 congeners with those of the analogous nitrido trifluorides $N \equiv MF_3$ of group 6 (M = Cr, Mo, W)^[25] and group 9 (Co, Rh, Ir)^[26] transition metals which have been studied previously. To the best of our knowledge, N≡Fe^{VI}F₃ is the first experimentally verified neutral, nitrido iron(VI) complex. In addition, we have evidence for the formation of $NM^{VII}F_4$ (M = Ru, Os).



For an electronic metal d^2 configuration of these $N \equiv MF_3$ compounds in C_{3v} symmetry Hund's rule predict that two parallel spin electrons occupy the degenerate $M(d_{xy,x^2-y^2})$ orbitals of e-type symmetry resulting in a non-degenerate highspin 3A_2 ground state. Although this 3A_2 state is not Jahn—Teller (JT) active, an electronic e^2 configuration can generally lead to a Jahn—Teller distorted ground state as a result of a strong pseudo-Jahn—Teller (PJT) mixing of two excited singlet electronic states. $^{[27]}$ This is because an electronic e^2 configuration in C_{3v} symmetry, in addition to the 3A_2 state, is generally associated with two electronic singlet states 1A_1 and 1E . These electronic states are reminiscent of the well-known singlet excited states of molecular oxygen. $^{[28]}$

It has been noted that the JT stabilization energy of the excited ¹E state is usually much weaker than the PJT stabilization resulting from mixing of the two excited ¹A₁ and ¹E states. The stabilization energy of this PJT interaction can be so large that the lower of these excited states crosses the ³A₂ potential energy surfaces and become the distorted global minimum configuration. ^[27,31] We observed such a "hidden" PJT distortion for N=RuF₃ and N=OsF₃ but not for NFe=F₃. Note that this distortion is also associated with a PJT-induced triplet-singlet spin crossover. ^[27a]

Results

Vibrational wavenumbers of group 8 nitrido trifluorides $\text{NM}^{\text{VI}}\text{F}_{\text{3}}$ and tetrafluorides $\text{NM}^{\text{VII}}\text{F}_{\text{4}}$

The IR spectra of the novel group 8 metal nitrido trifluorides, $N \equiv MF_3$ (M = Fe, Ru, Os) were recorded from the products obtained from laser-ablated free metal atoms with NF_3 seeded in a 1:1000 excess of neon after their deposition at 5 K on a gold-plated copper mirror (for experimental details see the Supporting Information). According to density functional theory calculations, the direct insertion of the metal atoms into an F-N bond of NF_3 to yield F_2N-MF , and the subsequent fluorine migration from nitrogen to the metal center to $FN=MF_2$ is highly exothermic for all three metals (Figure 1, Table S2).

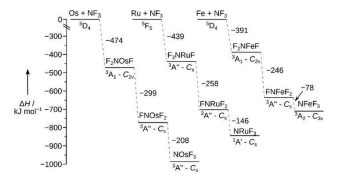


Figure 1. Stationary points on the reaction coordinate obtained at the BP86 level of theory for the formation of the nitrido metal complexes $N \equiv MF_3$ starting from the free metal atoms M and NF_3 (C_{3v} $^{-1}A_1$). See Table S2 for more details.

The rearrangement of the fluorimido complexes to the hexavalent nitrido trifluorides $N \equiv MF_3$ is found to be considerably exothermic for osmium ($-208 \text{ kJ} \, \text{mol}^{-1}$), ruthenium ($-146 \text{ kJ} \, \text{mol}^{-1}$), and iron ($-78 \text{ kJ} \, \text{mol}^{-1}$) at the BP86/def2-QZVP^[32] level of theory (details see the Supporting Information). Experimental IR spectra are shown from the deposits obtained in solid neon for the iron (Figures 2 and S1), ruthenium (Figures 3 and S2), and the osmium experiments (Figures 4 and S3), respectively. Experimental band positions are compared

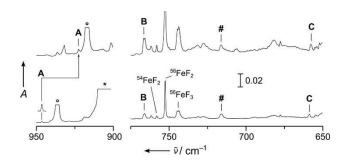


Figure 2. IR absorption spectra obtained from co-deposition of laser-ablated iron with 0.1 % $^{14}\text{NF}_3$ (bottom) and 0.1 % $^{15}\text{NF}_3$ (top) in solid Ne. Bands labeled with **A**, **B** and **C** are assigned to NFeF $_3$ (Table 1). Band **A** is enhanced by a factor of five. Known bands of binary iron fluorides $^{[29]}$ are labeled, and an unassigned band showing no $^{14/15}\text{N}$ isotopic shift is labeled with a hash mark. The bands associated with NF $_2$ and NF $_3$ are marked with circles and asterisks, respectively. $^{[30]}$ For more details, see Figure S1.

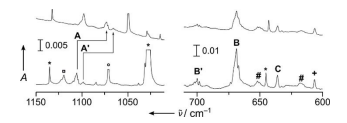


Figure 3. IR absorption spectra obtained from co-deposition of laser ablated ruthenium with 0.1% 14 NF $_3$ (bottom), and 15 NF $_3$ (top) in solid Ne, respectively. Bands labeled **A–C** are attributed to NRuF $_3$ and **A**′ and **B**′ are due to NRuF $_4$. Unknown bands are labeled by a pound and a plus sign, respectively. The bands associated with 14 NF, 14 NF $_2$ and 14 NF $_3$ are marked with squares, circles, and asterisks, respectively. [30] For more details, see Figure S2.

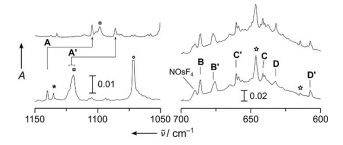


Figure 4. IR absorption spectra of laser ablated osmium co-deposited with 0.1 % 14 NF $_3$ in solid Ne (bottom), with 0.1 % 15 NF $_3$ in Ne (top). Bands labeled **A–D** are attributed to NOsF $_3$ (1 A') and **A'–D'** to NOsF $_3$ (3 A"). The bands marked with a pentagram sign are binary osmium fluorides. The bands associated with NF, NF $_2$ and NF $_3$ are marked with squares, circles and asterisks, respectively. [30] For more details see Figure S3.



with predicted ones from quantum-chemical calculations in Tables 1 and S4 (for a detailed band assignment refer to the Supporting Information). The formation of molecular NFeF₃ (C_{3v}) is clearly proved by the assignment of all its stretching vibrations marked **A** (ν (NFe): 946.4 cm⁻¹), **B** (ν_{as} (FeF₃): 766.8 cm⁻¹), and **C** (ν_s (FeF₃): 658.8 cm⁻¹) in Figure 2 (Table 1). Bands at 743.6/744.7, 752.6 and 785.1 cm⁻¹ were assigned to the known molecular binary iron fluorides ⁵⁶FeF₃, ⁵⁶FeF₂ and ⁵⁴FeF₂, respectively. ^[29] Their high intensity and the high yield of these binary fluorides compared to the NFeF₃ product bands indicate the lower stability of NFeF₃ under the harsh conditions of the laser ablation process. The spectra recorded in the ruthenium experiment (Figure 3), clearly revealed the presence of two different nitrido ruthenium complexes, finally assigned to NRuF₃ (C_s) and NRuF₄ (C_{4v}). The characteristic Ru \equiv N stretching bands of NRuF₃ (C_s) and NRuF₄ (C_{4v}) are labeled **A** (1105.4 cm⁻¹, NRuF₃) and A' (1098.5 cm⁻¹, NRuF₄) in Figure 3. The RuF₃ stretching modes of C_s symmetric NRuF₃ split into three modes. The strong antisymmetric F–Ru–F appears at 668.5 cm⁻¹ (labeled B in Figure 3) and likely overlaps with the nearby weaker F'-Ru band. The symmetric F-Ru-F mode is attributed to the band labeled **C** in Figure 3 at 635.8 cm⁻¹ (Table 1).

Table 1. Calculated and experimental vibrational wavenumbers (ν (^{14}N) in cm $^{-1}$) and $^{14/15}$ N isotopic shifts ($\Delta \nu$ in parentheses) for NFeF $_3$, NRuF $_4$, NOsF₃ and NOsF₄.

| Exp. ^[a] | CCSD (T) ^[b] | Assignment |
|---|-----------------------------|--|
| | | 3 |
| NFeF ₃ (C _{3v} , ³ A ₂) ^[c] | ((of | |
| 946.4 (-23.7) ^[d] | 1028 (-26) ^[e,f] | NFe str., a ₁ |
| 766.8/766.7 (0) | 737 (0) ^[f] | FeF₃ str., e |
| 658.8 (-1.1) | 689 (-2) ^[f] | FeF ₃ str., a ₁ |
| $N^{102}RuF_3 (C_s, {}^1A')^{[g]}$ | | |
| 1105.4 (-32.7) | 1085 (-32) | NRu str., a' |
| _ ^[h] | 682 (0) | F'-Ru str., a' |
| 668.5 (0) | 678 (0) | antisym. F–Ru-F str., a" |
| 635.8 (0) | 649 (0) | sym. F–Ru-F str., a' |
| $N^{102}RuF_4 (C_{4v}, {}^2B_1)^{[g]}$ | | |
| 1098.5 (-32.5) | 1080 (-32) | NRu str., a ₁ |
| 700.1 (0) | 711 (0) | RuF₄ stretch, e |
| _[h] | 681 (0) | RuF ₄ stretch, a ₁ |
| _ ^(j) | 598 (0) | RuF ₄ stretch, b ₂ |
| NOsF ₃ (C _s , ¹ A') | | |
| 1140.1 (-35.5) | 1152 (-36) | NOs str., a' |
| 686.0/686.6 (0) | 689 (0) | OsF ₂ sym. str., a' |
| 641.3/640.0 (0) | 664 (0) | OsF' sym. str., a' |
| 632.3 (0) | 652 (0) | OsF ₂ antisym. str., a" |
| NOsF ₃ (Cs, ³ A") | | · |
| 1086.0 (-) ^[i] | 1095 (-36) ^[i] | ¹⁵ NOs str, a' |
| 675.8/677.0 | 675 (0) | OsF ₂ antisym. str., a" |
| 660.5/658.9 (0) | 668 (0) | OsF ₂ sym. str., a' |
| 607.4 (0.0) | 614 (0) | OsF' sym. str., a' |
| NOsF ₄ (C_{4v} , ${}^{2}B_{1}$) | | |
| _ ^[k] | 1145 (-36) | NOs str., a ₁ |
| _ ^[k] | 706 (0) | OsF ₄ stretch, a ₁ |
| 689.9 (0) | 693 (0) | OsF ₄ stretch, e |
| _0 | 635 (0) | OsF ₄ stretch, b ₂ |
| | | |

[a] Neon matrix; matrix sites are separated by a slash. [b] Intensities from DFT calculations available in Table S4. [c] M06-L/def2-QZVP: 785 a_1 (-11) [12], 703 e (0) [200], 617 a₁ (-1) [40]. [d] ^{14/15}N isotopic ratio: 1.0256. [e] ¹⁴ ¹⁵N isotopic ratio: 1.0257. [f] NEVPT2/aug-cc-pwCVTZ-DK. [g] For the experimentally observed Ru isotope splitting see Tables S5-S7 and Figures S4 and S5). [h] Band is likely hidden by the stronger antisymmetric F-Ru-F stretching mode (a"). [i] $v(^{15}N-Os)$ in cm⁻¹, see text. [j] Not IR active. [k] Too weak or overlapped.

For NRuF₄ only the strongest RuF₄ stretching band, the degenerate e-type mode could safely be assigned to the band labeled B' in Figure 3 centered at 700.0 cm⁻¹.

In the spectra obtained from the reaction of osmium atoms with isotopic labeled ¹⁵NF₃ two Os=N stretching bands appeared at 1104.6 and 1086 cm⁻¹, which are labeled A and A', respectively, in Figure 4, and which are finally assigned to different "spinisomers" of NOsF₃ in near-degenerate singlet ¹A' and triplet ³A" electronic states (Table 1). In the ¹⁴NF₃ experiment A is observed at 1140 cm⁻¹ (Figure 4), while A' is overlapped by a stronger band due to the ¹⁴NF radical at 1120.8 cm⁻¹. [30b] All three Os—F stretching bands of singlet NOsF₃ (¹A') are assigned (Table 1) and labeled **B** $(\nu_s(OsF_2): 686.0 \text{ cm}^{-1}), \ C \ (\nu(OsF'): 641.3 \text{ cm}^{-1}), \ \text{and} \ D \ (\nu_{as}(OsF_2): 686.0 \text{ cm}^{-1})$ 632.3 cm⁻¹) in Figure 4, respectively. Bands labeled B', C' and D' at 675.8 cm⁻¹, 660.5 cm⁻¹ and 607.4 cm⁻¹, respectively, are assigned to the three Os-F stretching modes of triplet NOsF₃ (³A", Table 1). Finally, a band at 689.6 cm⁻¹ in Figure 4 is tentatively assigned to the strongest vibrational mode of $NOsF_4$ (C_{4v} , Table 1). The tetrafluorides $N \equiv MF_4$ (M = Ru, Os) are likely formed by the exothermic addition of a fluorine atom to $N=MF_3$ (Table S2).

Pseudo-Jahn-Teller distortion of molecular group 8 nitrido fluorides NM^{VI}F₃

The group 8 nitrido fluorides $NM^{VI}F_3$ adopt metal d^2 configurations, for which Hund's rule predicts a high-spin ³A₂ ground state in an undistorted C_{3v} symmetry and two parallel spin electrons in the twofold degenerate $e(d_{xy,x^2-y^2})$ -orbital $(|e_{\varepsilon}\uparrow;e_{\theta}\uparrow\rangle)$, labeled 9e for NFeF₃ in the Supporting Information Figure S6. Three e^2 terms (four states) can be formed, 3A_2 ($|e_{\epsilon}\uparrow;e_{\theta}\uparrow\rangle$), 1A_1 ($\sqrt{}$ $^{1}/_{2}$ [$|e_{\varepsilon}\uparrow;e_{\varepsilon}\downarrow\rangle+|e_{\theta}\uparrow;e_{\theta}\downarrow\rangle$]), $^{1}E_{\theta}$ ($\sqrt{^{1}}/_{2}$ [$|e_{\varepsilon}\uparrow;e_{\varepsilon}\downarrow\rangle-|e_{\theta}\uparrow;e_{\theta}\downarrow\rangle$]) and $^{1}E_{\varepsilon}$ $(\sqrt{1/2} \ [|e_0\uparrow;e_{\epsilon}\downarrow\rangle + |e_0\downarrow;e_{\epsilon}\uparrow\rangle])$. Due to the nondegenerate nature and totally symmetric charge distribution of the ³A₂ state no Jahn-Teller distortion is expected. [31] Other distributions of the electrons, as outlined above, result in configurations with lower spin and the absence of low-lying triplet excited states rule out obvious ground state pseudo-Jahn-Teller distortions.

Nevertheless, as shown in Figure 5 and in agreement with experimental vibrational assignments, all four NMF₃ species possess surprisingly different structures and the C_{3v} symmetric ground state was only verified for NFeF3. In case of NRuF3, extensive CCSD(T)/CBS calculations (Table S10) find the high symmetric ³A₂ is just about 5 kJ mol⁻¹ higher than the distorted ¹A' ground state. According to our experimental data, NOsF₃ features two quasi-degenerate, distorted structures in ¹A' and ³A" electronic states, separated by only $\Delta E_{T-S} = -1.3 \text{ kJ mol}^{-1}$ (CCSD(T)/CBS, Table S11).

To elucidate these findings, adiabatic potential energy surface (APES) scans were carried out using state-averaged complete active space self-consistent field calculations by distributing eight electrons in the eight molecular orbits formed by the metal (n-1)d and N(2p) orbitals (SA-CASSCF(8,8)) with subsequent NEVPT2 treatment to recover dynamic correlation. Shown in Figure 6a-c are cross sections along a distortion coordinate (D) that connects the two stationary points of the

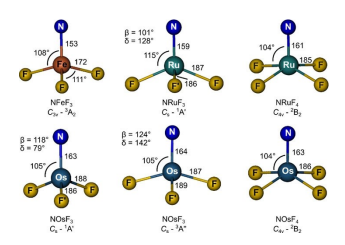


Figure 5. Ground-state structures of NMF₃ and NMF₄ calculated at the CCSD (T)/aVTZ (M = Ru, Os, M: aVTZ-PP) and the NEVPT2/aVTZ-DK (NFeF₃, Fe: awCVTZ-DK) levels of theory. Bond lengths are given in pm and angles in degrees. β denotes the N–M–F' and δ the F–M–F angle for structures with C_s

 1 A' surface, at D = -1 and 1, respectively, via the high-symmetry C_{3v} stationary point at D=0.

The distortions take place along one component of the lowest (NFeF₃, NRuF₃) or imaginary (NOsF₃) degenerate e normal mode in the high-symmetry C_{3v} configuration. Therefore, mainly bond angle distortions are involved, in particular the dihedral angle F'-M-N-F (α , Figure S9), and the valence angles N-M-F' (β , Figure 5), and N–M–F (γ). The sign of the distortion D in Figure 6 indicates a widening (positive) or closing (negative) of α . Differences in these angles and in the three nonequivalent bond distances between two localized stationary points in C_s symmetry were divided into equal incremental steps and used as intermediate internal coordinates in the APES calculation for each step (Tables S14-S17). In the case of Figure 6d the distortion in the positive direction was carried out using the NOsF₃ ³A" minimum structure at D=1. The graphs shown in Figure 6a–d represent the energies of the terms arising from the electronic e^2 configuration, as outlined above. They demonstrate the propensity of trigonal group 8 nitrido complexes in the oxidation state VI to be subject to a PJT distortion. Other trigonal systems displaying a (A+E) \otimes e Pseudo-Jahn-Teller effect (PJTE) that is "hidden" in excited states (h-PJTE) have already been described.[27a,31] The condition for a distorted ground state minimum structure caused by the h-PTJE is that the PJT stabilization energy of an excited state (E_{PJT}) is larger than the energy gap Δ_0 between the ground state in the highsymmetry configuration and the PJT active excited state ($E_{\rm PJT} > \Delta_{\rm O}$, see Figure 6, a-c).[27a] The global minimum of the APES of NFeF3 shown in Figure 6a is located at the high-symmetry point. The stationary points on the ¹A' (blue line) surface are a local minimum (D=1) and a first-order saddle point (D=-1) without surface crossings in between. Consistent with the experimental vibrational data the global minimum is the high symmetry configuration. The h-PJTE in the ¹E state is not strong enough to distort the high-

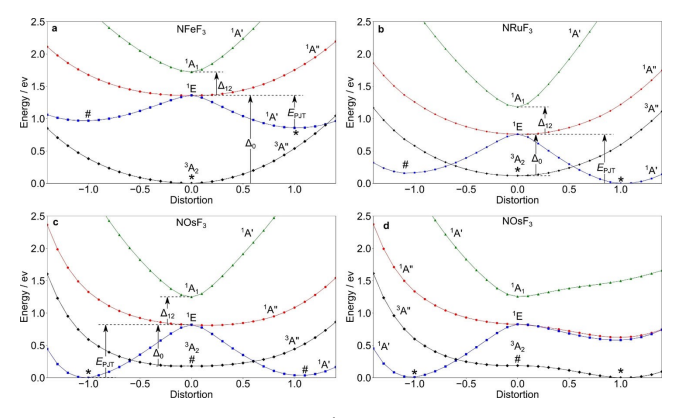


Figure 6. Cross section of the APES for the terms arising from the electronic e^2 configurations of a) NFeF₃, b) NRuF₃, c), d) NOsF₃ along the distortion coordinate (D) connecting stationary points located at D=1, 0 and -1, respectively, on the $^{1}A'$ (blue line) and the $^{3}A_{2}$ ($^{3}A''$, black line) surfaces. Minimum points are marked with an asterisk, and first-order saddle points with a hash mark. The PJT stabilization energy (E_{PJT}) of the lowest excited state, its excitation energy at C_{3v} symmetry (Δ_0), and the $^1E-^1A_1$ energy gap (Δ_{12}) are indicated in (a)–(c).



symmetry configuration. The PJT stabilization energy, $E_{\rm PJT}$, is about 0.39 eV and smaller than the $^{1}\text{E-}^{3}\text{A}_{2}$ energy gap $\Delta_{0}=1.36$ eV. The $^{1}\text{A}'$ minima, which features a ($|e_{0}\uparrow:e_{0}\downarrow\rangle$) electronic configuration, and the $^{3}\text{A}_{2}$ minima are separated by about 0.86 eV. The angular distortion from D=-1 to D=1 at the $^{1}\text{A}'$ surface extends from about $101^{\circ}-132^{\circ}$ (α), $100^{\circ}-120^{\circ}$ (β), and $112^{\circ}-104^{\circ}$ (γ).

The cross section of the APES of NRuF₃ along the distortion coordinate from D=0 to D=1 illustrated in Figure 6b shows that one of the components of the ¹E term is stabilized by the strong PJT coupling with the excited ¹A₁ state. It crosses the ³A₂ ground state of the undistorted high-symmetry configuration to produce the global minimum with a distorted structure. The triplet-singlet spin crossover is associated with an orbital disproportionation, [27a] because in the distorted structure the electrons are paired in one e_0 orbital ($|e_0\uparrow;e_0\downarrow\rangle$) instead of the symmetric distribution ($|e_e\uparrow;e_0\uparrow\rangle$) in the undistorted configuration. Accordingly, we find that $E_{P,\Pi}=0.76$ eV is larger than $\Delta_0=0.64$ eV. The high-spin ³A" state is higher in energy by only ~0.12 eV and it has an energy barrier of ~0.25 eV to the point of spin crossover with the low-spin ¹A' state.

Figures 6c and d exhibit four relevant low-lying stationary points on the ¹A' and ³A" APES of NOsF₃. The h-PJTE in this case produces a minimum with a distorted ${}^{1}A'$ structure at D=-1and accordingly, the orbital disproportionation and spin crossover leads to a ($|e_{\epsilon}\uparrow; e_{\epsilon}\downarrow\rangle$) configuration with $E_{PJT} = 0.82$ eV and $\Delta_0 = 0.60$ eV. Unlike the former two cases, the 3A_2 highsymmetry configuration of NOsF3 does not represent a minimum point, but a first order saddle point. Following the ϵ component of the imaginary e mode in Figure 6d we find – in accordance with the CCSD(T)/CBS results - an energetically quasi-degenerate distorted ³A" minimum that shows orbital disproportionation, but no spin crossover about 0.1 eV (or 0.7 kJ mol⁻¹) lower than the ¹A' state. The energy barrier of the spin crossover point is ~0.27 eV (CCSD(T)/VTZ-PP: 0.24 eV, Table S12), a significant barrier connecting both stationary points at the experimental cryogenic conditions. These findings support the observation of two different species in the experimental infrared spectra which correspond to species in different ¹A' and ³A" electronic states. We did not analyze the source of the distortion of the high-spin minimum (³A"). But, under the premise that PJTE is the only source for symmetry breaking of non-degenerate high-symmetry states, [27b,31] the source is most likely an interacting triplet ³E excited state.

Discussion

All metal specific bands showing a ^{14/15}N isotopic shift were successfully assigned. Bands due to binary fluorides are always present in experiments using IR laser ablation of metals in the presence of molecular fluorides as precursors. They are likely formed by recombination of metal atoms and atomic fluorine radicals formed by thermal or photolytic decomposition of the fluoride precursor in the hot plasma plume region or by the decomposition of metal fluoride product molecules. However, the very strong NF₃ precursor bands and comparatively weak NF and NF₂ bands in all spectra suggest that the formation of the NMF₃

title product can be attributed to the reaction of M and NF₃. Lower nitrido fluorides NMF or NMF₂ could in principle also be formed through the cleavage of a metal-fluorine bond or by the reaction of metal atoms with NF or NF₂, but have so far not been identified. ^[25,26,30c,33] The addition of fluorine to NMF and NMF₂ to yield NMF₃ and also the formation of NMF₄ for M=Ru and Os are calculated to be exothermic (Table S2).

As shown here, all the trigonal NMF₃ species possess two equilibrium configurations with different spin multiplicities, while those of NRuF₃ and NOsF₃ are close in energy. Such a magnetic and structural PJT induced bistability may also be possible for ligand-stabilized trigonal nitrido d² metal complexes. Such compounds are of interest for molecular switching, especially when symmetry breaking is involved (as for NFeF₃ and NRuF₃).^[34]

The different stationary structures that were obtained for the group 8 NMF₃ molecules shown in Figure 5 possess surprisingly different electronic configurations, as outlined above and summarized in Figure 7 (for molecular orbital plots, see Figures S6 and S8). The different 1 A' electronic ground states of NRuF₃ and NOsF₃ arise from the pairing of two unpaired electrons in different orbitals, which are associated with two different structural distortions. The HOMO of NRuF₃ (1 A') is of a" symmetry, which is consistent with a widening of the F–M–F angle, whereas the HOMO of NOsF₃ (1 A') is of a' symmetry, which shows a reduction in the F–M–F angle bisected by the σ plane in $C_{\rm s}$ symmetry (Figure 5). The d¹ metal configuration for the heptavalent tetrafluorides NRu $^{\rm NIF}_4$ and NOs $^{\rm NIF}_4$ ($C_{\rm 4v}$) give rise to a 2 B₂ electronic ground state (see Figure S7 for the singly occupied MO).

The effective bond orders^[35] (EBOs) for NOsF₃, NRuF₃ and NFeF₃ are 2.8, 2.7 and 2.2, respectively, which in fact corresponds to triple bonds for all these M—N bonds. The computed M—N bond lengths for the novel nitrido compounds (153 pm (FeN), 159 pm (RuN), 163–164 pm (OsN), Figure 7) are close to our published triple bond additive covalent radii: 156 pm (FeN), 157 pm (RuN) and 163 pm (OsN), [36] and also the experimental N—M stretching frequencies (Table 1) support the presence of strong M \equiv N triple bonds in the novel hexavalent nitrido complexes NM^{VI}F₃. We note that the experimental ν (Fe \equiv N) frequency of NFe^{VI}F₃ of 946 cm $^{-1}$ (Table 1) is not well reproduced by calculations at DFT or CCSD(T) levels (Table S3) and is also

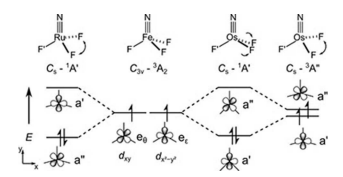


Figure 7. Comparison of the different d^2 electron configurations of the NMF₃ species (M = Fe, Ru, Os). The metal centered a' and a"-MOs are dominantly $M(d_{x^2-y^2})$ and $M(d_{xy})$ atomic orbitals, respectively.



overestimated by the more sophisticated NEVPT2 multi-reference approach ($\nu(\text{Fe}=\text{N})=1027~\text{cm}^{-1}$, Table S3). On the other hand, its comparison with experimental Fe=N stretching frequencies for pseudo-tetrahedral NIVFeL3 complexes, previously reported at 1008 ([FeIV(N)(TIMENIMES)]+),[24c] 1028 ([FeIV(N)(PhB('Bulm)_3]),[24b] and 1034 cm⁻¹ ([FeIV(N)-(PhB(CH_2P'Pr_2)_3],[49] Table S1) suggests that an increase in the iron oxidation state beyond V does not necessarily lead to a stronger Fe=N bond.

Table 2 shows experimental M–N stretching frequencies of molecular NMF₃ species formed by the reaction of NF₃ with laserablated transition metals. For the d⁰ configurations of all group 4 and group 6 nitrido trifluorides the ideal pseudo-tetrahedral $C_{3\nu}$ symmetric arrangement was experimentally verified, since there are no electrons in the nonbonding $e(d_{xy,x^2-y^2})$ orbitals that could cause distortions.^[25,33] The e^3 configuration of NRhF₃ and NIrF₃ leads to Jahn–Teller distorted spin doublet ground states in C_s symmetry.^[26] So far, no experimental data are available for the group 10 derivatives, and for the group 11 analogues only the initial metal insertion products F₂N–M^{II}F were detected after matrix deposition (irradiation of F₂NCuF led to rearrangement to metastable FN–CuF₂).^[30c]

The M-N stretching normal mode of the terminally bond nitrogen ligands of the nitrido trifluorides can regarded to be a good approximation as an almost pure and uncoupled metalnitrogen stretching mode that can be used as a measure of the M–N bond strength. The NM^{IV}F₃ derivatives of the group 4 metals possess a singly bonded triplet nitrene (3N-) ligand, since the ligand cannot oxidize the d⁰ metal center any further. The two unpaired electrons in the N(2p) orbitals are reported to be involved in weak degenerate π bonding interactions for M=Ti \gg Zr, Hf.[33] In contrast, the group 6, 8 and 9 NMVF₃ molecules show a N \equiv M triple bond with one σ and two π bonds to the terminal nitrido (N³-) ligand. The strength and overlap of these bonds increases going down the groups likely due to an improved $M(\pi d)$ – $N(\pi p)$ orbital overlap as a result of an increasing relativistic expansion^[37] of the 4d and 5d orbitals and the absence of metal core/ligand repulsion proposed in first-row transition metal compounds.[38] The general trend of increasing N-M bond strength moving along the rows culminates in the highest observed M-N stretching frequency for NIrF₃. Unexpectedly, this trend does not apply to NFeF₃ which shows a lower M-N stretching frequency than the group 6 homologue (M = Cr). The lower stability of high-valent first row late transition metals is well known. $^{\![1a,39]}$ In the series of 3d $NM^{VI}F_3$ compounds, for $M\!=\!Fe$ it seems we have reached the limit of stability. NCo^{VI}F₃ is not a stable compound and only FNCoF₂ has been observed experimentally.^[26] For the 4d element Rh it was found that the rearrangement of the fluoro nitrene complex $FNRhF_2$ into $N = Rh^{VI}F_3$ is only slightly exothermic ($\Delta H^0 = -12 \text{ kJ mol}^{-1}$, CCSD(T)), which enables the observation of both rearrangement products. [26]

Within the atoms in molecules (AIM) scheme^[40] the partial negative charge at the nitrido ligand in NMVF₃ increases from M= Fe to Os (Tables S9 and S13), which indicates a decreasing electron withdrawing effect of the MVF₃ fragment within this group. For M=Fe and Ru the negative charge at the nitrogen atom also decreases from NMF₂ (M=Fe: -0.35, Ru: -0.40) to NMF₃ (M=Fe: -0.25, Ru: -0.35), while for M=Os it remains unchanged (NOsF₂: -0.50, NOsF₃: -0.49). As expected, fluorination of NMF₃ further decreases the atomic charge of the nitrido ligand in NMF₄ (M = Ru: -0.25, Os: -0.40, Table S9). The high oxidation potential of Fe^{VI} in NFeF₃ leads to relatively high σ^* and π^* occupation numbers (0.2 and 0.3 electrons, respectively; Figure S6). These indicates a weakened covalent N-Fe bond, for which the formal N³⁻ nitride notations seems to be a very poor approximation. The occupation of formally antibonding MOs also indicates an oxidation, and thus the onset of a redox non-innocent behavior of the nitrido ligand.

Conclusion

The nitrido complexes NFeF₃, NRuF₄, NOsF₃ (¹A'), NOsF₃ (³A"), and NOsF₄ were shown to be formed by the reaction of free group 8 metal atoms with NF₃ and established by their characteristic IR spectra recorded in solid neon matrices. Their assignment is supported by observed 14/15N isotope shifts and quantum-chemical predictions. All stretching fundamentals of the NMVF₃ complexes were confidently assigned. For the C_{4v} symmetric NRuF₄ two distinct bands were confidently assigned, whereas for NOsF₄ only the strongest band was tentatively assigned. Based on the joint experimental IR and quantum-chemical analysis the half-filled e^2 configuration of NFeF₃ can be assigned to an undistorted C_{3v} structure in a non-degenerate ³A₂ electronic ground state. NFeF₃ features an unprecedented low Fe=N triple-bond frequency of 946.7 (14 N \equiv Fe) and 922.7 cm $^{-1}$ (15 N \equiv Fe). The heavier group 8 NMF $_3$ homologues are subject to symmetry lowering and spin-crossover caused by a pseudo Jahn-Teller effect "hidden" in the excited states. While the electronic ground state of NRuF₃ is a structurally distorted singlet ¹A' state (C_s symmetry), for molecular NOsF₃ two coexisting distorted C_s structures with high-spin and low-spin d² configurations (magnetic bistability) were detected at 5 K in solid neon. To the best of our knowledge, apart from $O_2Fe(\eta^2-O_2)$, [3-5] no other neutral Fe^{VI} complexes or molecular neutral complexes of Ru^{VII} have yet been reported, and after OsOF₅, [1f] NOsF₄ is the second known monomeric Os^{VII} compound.

| Row | Group 4 ^[a] | Group 6 ^[b] | Group 8 ^[c] | Group 9 ^[d] | Group 11 ^[30c] |
|------------|--|---|--|--|---|
| 3 <i>d</i> | 596.7 (Ti, C _{3v} , ³ A ₁) | 1015 (Cr, C _{3v} , ¹ A ₁) | 946.4 (Fe, C _{3v} , ³ A ₂) | FN=CoF ₂ only | F ₂ N-CuF, FN=CuF ₂ |
| 4 <i>d</i> | 553.1 (Zr, C_{3v} , ${}^{3}A_{1}$) | 1075 (Mo, C_{3v} , ${}^{1}A_{1}$) | 1098.5 (Ru, C _s , ¹ A') | 1116.1 (Rh, <i>C_s</i> , ² A′) [Ne] 1112.5 [Rh, Ar] ^[e] | F₂N–AgF only |
| 5 <i>d</i> | 548.1 (Hf, C_{3v} , ${}^{3}A_{1}$) | 1091 (W, C _{3v} , ¹ A ₁) | 1140.1 (Os, C _s , ¹ A') 1086.0 (Os, C _s , ³ A") | 1150.4 (lr, C _s , ² A') [Ne] 1144.6 [lr, Ar] | F₂N—AuF only |



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Conflict of Interest

The authors declare no conflict of interest.

Keywords: ab initio calculations · high oxidation states · matrix isolation · N ligands · transition metals

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